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- (75) Inventors/Applicants (for US only): EMINI, Emilio, A. [US/US]; 126 East Lincoln Avenue, Rahway, NJ 07065-0907 (US). YOUIL, Rima [AU/US]; 126 East Lincoln Avenue, Rahway, NJ 07065-0907 (US). BETT, Andrew, J. [CA/US]; 126 East Lincoln Avenue, Rahway, NJ 07065-0907 (US). CHEN, Ling [US/US]; 126 East Lincoln Avenue, Rahway, NJ 07065-0907 (US). KASLOW, David, C. [US/US]; 126 East Lincoln Avenue, Rahway, NJ 07065-0907 (US). SHIVER, John, W. [US/US]; 126 East Lincoln Avenue, Rahway, NJ 07065-0907 (US). TONER, Timothy, J. [US/US]; 126 East Lincoln Avenue, Rahway, NJ 07065-0907 (US). CASIMIRO, Daniel, R. [PH/US]; 126 East Lincoln Avenue, Rahway, NJ 07065-0907 (US).
- (74) Common Representative: MERCK & CO., INC.; 126
 East Lincoln Avenue, Rahway, NJ 07065-0907 (US).

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(54) Title: ENHANCED FIRST GENERATION ADENOVIRUS VACCINES EXPRESSING CODON OPTIMIZED HIV1-GAG, POL, NEF AND MODIFICATIONS

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(57) Abstract: First generation adenoviral vectors and associated recombinant adenovirus-based HIV vaccines which show enhanced stability and growth properties and greater cellular-mediated immunity are described within this specification. These adenoviral vectors are utilized to generate and produce through cell culture various adenoviral-based HIV-1 vaccines which contain HIV-1 gag, HIV-1 pol and/or HIV-1 nef polynucleotide pharmaceutical products, and biologically relevant modifications thereof. These adenovirus vaccines, when directly introduced into living vertebrate tissue, preferably a mammalian host such as a human or a non-human mammal of commercial or domestic veterinary importance, express the HIV1- Gag, Pol and/or Nef protein or biologically modification thereof, inducing a cellular immune response which specifically recognizes HIV-1. The exemplified polynucleotides of the present invention are synthetic DNA molecules encoding HIV-1 Gag, encoding codon optimized HIV-1 Pol, derivatives of optimized HIV-1 Pol (including constructs wherein protease, reverse transcriptase, RNAse H and integrase activity of HIV-1 Pol is inactivated), HIV-1 Nef and derivatives of optimized HIV-1 Nef, including nef mutants which effect wild type characteristics of Nef, such as myristylation and down regulation of host CD4. The adenoviral vaccines of the present invention, when administered alone or in a combined modality regime, will offer a prophylactic advantage to previously uninfected individuals and/or provide a therapeutic effect by reducing viral load levels within an infected individual, thus prolonging the asymptomatic phase of HIV-1 infection.

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IPC(7) US CL	SSIFICATION OF SUBJECT MATTER : C12N 15/86 : 435/456				
	International Patent Classification (IPC) or to both r. DS SEARCHED	national classification and IPC			
Minimum documentation searched (classification system followed by classification symbols) U.S.: 424/205.1, 207.1, 227.1, 233.1; 435/69.1, 69.3, 173.3, 235.1, 320.1, 456; 530/23.72;					
Documentati	on searched other than minimum documentation to th	e extent that such documents are include	d in the fields searched		
	ata base consulted during the international search (nar continuation Sheet	ne of data base and, where practicable, s	earch terms used)		
	UMENTS CONSIDERED TO BE RELEVANT				
Category *	Citation of document, with indication, where ap		Relevant to claim No.		
X 	WO 96/39178 (ERTL et al.) 12 December 1996 (12 and claims 1 and 5.	2.12.1996), see page 5, 6,10, 12, 13	1-3, 8-11, 18		
Y	VID 4 040 000 4 70000		4, 5, 13-17, 29-32, 34, 35, 37		
X 	US 6,019,978 A (ERTL et al.) 1 February 2000, (01	1/02/2000), see columns 2, 7 and 8.	1-3, 8-11, 18		
Y			4, 5, 13-17, 29-32, 34, 35, 37		
X,P	US 6,287,571 β (ERTL et al.) 11 September 200 and claim 1.	01 (11/09/2001), see columns 2, 7, 8	1, 9, 18		
X	US 5,643,579A (HUNG et al.) 1 July 1997 (01/07/	1997), see examples 1, 2, 25 and 26.	1-3, 8, 9-11, 18		
Ÿ			4,5,13-17, 29-32, 34, 35, 37		
Y	WANG et al. The use of an E1-deleted, replication expressing the rabies virus glycoprotein for early virunal of Virology (March 1997) Vol. 71, No. 5,	accination of mice against rabies virus.	1-3, 9-11, 13-18		
Further	documents are listed in the continuation of Box C.	See patent family annex.			
* S	pecial categories of cited documents:	"T" later document published after the in priority date and not in conflict with			
	t defining the general state of the art which is not considered to ticular relevance	understand the principle or theory u "X" document of particular relevance; th	nderlying the invention		
"E" carlier ap	oplication or patent published on or after the international filing	considered novel or cannot be consisted when the document is taken alo	dered to involve an inventive		
	t which may throw doubts on priority claim(s) or which is cited is the publication date of another citation or other special reason fied)	"Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such			
"O" documen	combination being obvious to a person skilled in the art document referring to an oral disclosure, use, exhibition or other means "&" document member of the same patent family				
Date of the a	Date of the actual completion of the international search Of February 2002 (06.02.2002) Date of mailing of the international search report 19 AUG 2002				
	ailing address of the ISA/US	Authorized officer	1,10		
Con Box	omissioner of Patents and Trademarks PCT chington, D.C. 20231	Ulrike Winkler, Ph.D.			
	o. (703)305-3230	Telephone No. 703-308-0196			

Form PCT/ISA/210 (second sheet) (July 1998)

International application No.

PCT/US01/28861

ategory *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No
Υ .	NATUK et al. Immunogenicity of recombinant human adenovirus -human immunodeficiency virus vaccines in chimpanzees. Aids Research and Human Retroviruses (1993) Vol. 9, No. 5, pp395-404, see material and methods.	1, 9, 29-32
Y	PREVEC et al. Immune response to HIV-1 gag antigens induced by recombinant adenovirus vectors in mice and rhesus macaque monkeys. Journal of Acquired Immune Deficincy Syndrome. (1991) Vol. 4, No. 6 pp. 568-76, see abstract.	1, 9, 29-32
Y	LORI et al. Rapid protection against human immunodeficiency virus type 1 (HIV-1) replication mediated by high efficiency non-retroviral delivery of genes interfering with HIV-1 tat and gag. Gene Therapy (1994) Vol. 1, No. 1, pp. 27-31, see abstract.	1, 9
Y	PFARR et al. Differential effects of polyadenylation regions on gene expression in mammalian cells. DNA (1986) Vol. 5, No. 2, pp.115-22, see abstract.	16
Y	NATUK et al. Adenovirus vectored vaccine. Developmental Biological Standards (1994) Vol. 82, pp. 71-77, see abstract.	. 1,9
	,	
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International application No.

PCT/US01/28861

Box I Observations where certain claims were found unsearchable (Continuation of Item 1 of first sheet)		
This international report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:		
Claim Nos.: because they relate to subject matter not required to be searched by this Authority, namely:		
2. Claim Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:		
3. Claim Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).		
Box II Observations where unity of invention is lacking (Continuation of Item 2 of first sheet)		
This International Searching Authority found multiple inventions in this international application, as follows: Please See Continuation Sheet		
As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.		
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.		
As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:		
4. No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.: 1-5, 8-11, 13-18, 29-32, 34, 35, 37		
Remark on Protest The additional search fees were accompanied by the applicant's protest.		
No protest accompanied the payment of additional search fees.		

International application No.

PCT/US01/28861

BOX II. OBSERVATIONS WHERE UNITY OF INVENTION IS LACKING

This application contains the following inventions or groups of inventions which are not so linked as to form a single general inventive concept under PCT Rule 13.1. In order for all inventions to be examined, the appropriate additional examination fees must be paid.

Group	Claims	
1	1-5, 8-11, 13-18, 29, 30, 31, 32, 34, 35, 37	The claims are directed to an adenoviral vector that is at least partially deleted of $\Delta E1$, the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Gag protein (SEQ ID NO: 29) inserted in the parallel orientation of E1. In addition the vector contains a promoter and a polyadenylation signal.
2	6, 7, 36	The claims are directed to an adenoviral vector that is at least partially deleted of $\Delta E1$ and $\Delta E3$, the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Gag protein (SEQ ID NO: 29).
3	12, 33	The claims are directed to an adenoviral vector that is at least partially deleted of $\Delta E1$, the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV protein inserted in the antiparallel orientation of E1.
4	19-23, 38-42	The claims are directed to a method of making and harvesting of a recombinant adenoviral particle that contains a gene encoding an HIV Gag protein.
5	24, 27, 28, 43, 46, 47	The claim is directed to a method of generating a cellular mediated immune response to HIV Gag protein with the recombinant adenoviral particle.
6	25, 26, 44, 45	The claim is directed to a method of generating a cellular mediated immune response to HIV Gag protein with the recombinant adenoviral particle in addition to administering a DNA plasmid vaccine.
7	48-51, 53, 54, 56	The claims are directed to an adenoviral vector that is at least partially deleted of $\Delta E1$, the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Pol protein (SEQ ID NO: 1) inserted in the parallel orientation of E1.
8	48-51, 53, 54, 56	The claims are directed to an adenoviral vector that is at least partially deleted of $\Delta E1$, the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Pol protein (SEQ ID NO: 5) inserted in the parallel orientation of E1.
9	48-51, 53, 54, 56	The claims are directed to an adenoviral vector that is at least partially deleted of $\Delta E1$, the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Pol protein (SEQ ID NO: 7) inserted in the parallel orientation of E1.
10	52	The claim is directed to an adenoviral vector that is at least partially deleted of $\Delta E1$, the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Pol protein (SEQ ID NO: 1) inserted in the antiparallel orientation of E1.
11	52	The claim is directed to an adenoviral vector that is at least partially deleted of $\Delta E1$, the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Pol protein (SEQ ID NO: 5) inserted in the antiparallel orientation of E1.
12	52	The claim is directed to an adenoviral vector that is at least partially deleted of $\Delta E1$, the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Pol protein (SEQ ID NO: 7) inserted in the antiparallel orientation of E1.
13	55	The claim is directed to an adenoviral vector that is at least partially deleted of $\Delta E1$

Form PCT/ISA/210 (second sheet) (July 1998)

International application No.

PCT/US01/28861

		and $\Delta E3$, the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Pol protein (SEQ ID NO: 1) inserted in E1.
14	55	The claim is directed to an adenoviral vector that is at least partially deleted of $\Delta E1$ and $\Delta E3$, the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Pol protein (SEQ ID NO: 5) inserted in E1.
15	55	The claim is directed to an adenoviral vector that is at least partially deleted of $\Delta E1$ and $\Delta E3$, the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Pol protein (SEQ ID NO: 7) inserted in E1.
16	57-61	The claims are directed to a method of making and harvesting of a recombinant adenoviral particle that contains a gene encoding an HIV Pol protein.
17	62, 65, 66	The claim is directed to a method of generating a cellular mediated immune response to HIV Pol protein with the recombinant adenoviral particle.
18	63, 64	The claim is directed to a method of generating a cellular mediated immune response to HIV Pol protein with the recombinant adenoviral particle in addition to administering a DNA plasmid vaccine.
19	67-70, 72, 73, 75	The claims are directed to an adenoviral vector that is at least partially deleted of $\Delta E1$, the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Nef protein (SEQ ID NO: 9) inserted in the parallel orientation of E1.
20	67-70, 72, 73, 75	The claims are directed to an adenoviral vector that is at least partially deleted of $\Delta E1$, the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Nef protein (SEQ ID NO: 11) inserted in the parallel orientation of E1.
21	67-70, 72, 73, 75	The claims are directed to an adenoviral vector that is at least partially deleted of $\Delta E1$, the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Nef protein (SEQ ID NO: 13) inserted in the parallel orientation of E1.
22	67-70, 72, 73, 75	The claims are directed to an adenoviral vector that is at least partially deleted of $\Delta E1$, the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Nef protein (SEQ ID NO: 15) inserted in the parallel orientation of E1.
23	71	The claim is directed to an adenoviral vector that is at least partially deleted of $\Delta E1$, the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Nef protein (SEQ ID NO: 9) inserted in the antiparallel orientation of E1.
24	71	The claim is directed to an adenoviral vector that is at least partially deleted of $\Delta E1$, the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Nef protein (SEQ ID NO: 11) inserted in the antiparallel orientation of E1.
25	71	The claim is directed to an adenoviral vector that is at least partially deleted of $\Delta E1$, the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Nef protein (SEQ ID NO: 13) inserted in the antiparallel orientation of E1.
26	71	The claim is directed to an adenoviral vector that is at least partially deleted of $\Delta E1$, the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Nef protein (SEQ ID NO: 15) inserted in the antiparallel orientation of E1.
27	74	The claim is directed to an adenoviral vector that is at least partially deleted of $\Delta E1$ and $\Delta E3$, the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Nef protein (SEQ ID NO: 9) inserted in E1.
28	74	The claim is directed to an adenoviral vector that is at least partially deleted of $\Delta E1$ and $\Delta E3$, the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Nef protein (SEQ ID NO: 11) inserted in E1.
29	74	The claim is directed to an adenoviral vector that is at least partially deleted of ΔE_1

International application No.

PCT/US01/28861

		adenovirus genome, and a gene which encodes an HIV Nef protein (SEQ ID NO: 13) inserted in E1.	
30	74	The claim is directed to an adenoviral vector that is at least partially deleted of $\Delta E1$ and $\Delta E3$, the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Nef protein (SEQ ID NO: 15) inserted in E1.	
31	76-80	The claims are directed to a method of making and harvesting of a recombinant adenoviral particle that contains a gene encoding an HIV Nef protein.	
32	81, 84, 85	The claims are directed to a method of generating a cellular mediated immune response to HIV Nef with the recombinant adenoviral particle.	
33	82, 83	The claims are directed to a method of generating a cellular mediated immune response to HIV Nef with the recombinant adenoviral particle in addition to administering a DNA plasmid vaccine.	
34	86a	The claim is drawn to a multivalent vaccine wherein gag, pol and nef are expressed from three individual vectors.	
35	86b, 88, 89	The claims are drawn to a multivalent vaccine wherein gag, pol and nef are expressed from one individual vectors.	
36	86c, 88	The claims are drawn to a multivalent vaccine wherein gag, pol and nef are expressed from two individual vectors, one expressing nef-pol fusion and one expressing gag.	
37	86d, 87, 88	The claims are drawn to a multivalent vaccine wherein gag, pol and nef are expressed from two individual vectors, one expressing gag-pol fusion and one expressing nef.	
38	86e, 88	The claims are drawn to a multivalent vaccine wherein gag, pol and nef are expressed from two individual vectors, one expressing nef-gag fusion and one expressing pol.	
39	86f, 88	The claims are drawn to a multivalent vaccine wherein gag, pol and nef are expressed from a single vectors as a fusion protein.	
40	86g, 88	The claims are drawn to a multivalent vaccine wherein gag and pol are expressed from two individual vectors.	
41	86h, 88, 89	The claims are drawn to a multivalent vaccine wherein gag and pol are expressed individually from one vector.	
42	86i, 88	The claims are drawn to a multivalent vaccine wherein pol and nef are expressed from two individual vectors.	
43	86j, 88, 89	The claims are drawn to a multivalent vaccine wherein <i>pol</i> and <i>nef</i> are expressed from individually from one vector.	
44	86k, 88	The claims are drawn to a multivalent vaccine wherein nef and gag are expressed individually from one vector.	
45	861, 88, 89	The claims are drawn to a multivalent vaccine wherein nef and gag are expressed individually from one vector.	
46	86m, 88	The claims are drawn to a multivalent vaccine wherein gag and pol are expressed as a fusion protein from one vector.	
47	86n, 88	The claims are drawn to a multivalent vaccine wherein <i>pol</i> and <i>nef</i> are expressed as a fusion protein from one vector.	
48	860, 88	The claims are drawn to a multivalent vaccine wherein nef and gag are expressed as a fusion protein from one vector.	

The inventions listed as Groups 1-48 do not relate to a single general inventive concept under PCT Rule 13.1 because, under PCT Rule 13.2, they lack the same or corresponding special technical features for the following reasons:

The technical feature linking groups 1-33 appears to be a recombinant adenoviral vector wherein the adenoviral vector is at least partially deleted in E1 but the vector may contain more deletions, the vector contains wild type sequences including packaging signals and a gene encoding a heterologous HIV protein or fragments thereof. Ertl et al. (WO 96/39178) disclose a recombinant adenoviral vector that is deleted in E1 and partially deleted in E3, the remainder of the adenoviral vector contains wild type sequences. The vector additionally contains an insertion of a heterologous protein which includes HIV proteins (see abstract and claims 1 and 5). Therefore, the technical feature linking the inventions of groups 1-45 does not constitute a special technical feature as defined by PCT Rule 13.2, as it does not define a contribution over the prior art.

The special technical feature of the following groups 1-3, 7-15, 19-30 and 34-48 is considered to be the combination of sequences that is disclosed in each group, see individual claim groupings above for the different sequences. The DNA disclosed in each group is made up of a different sequence having a different structure and different function.

International application No.

PCT/US01/28861

The special technical feature of group 4, 16 and 31 is considered to be a method of producing recombinant adenoviral particles. Each group contains different sequences hence the resulting particles would have different structures and functions associated with the particle.

The special technical feature of group 5, 17 and 32 is considered to be a method of producing a cellular mediated immune response to the heterologous protein encoded by the different adenoviral vectors. Each group contains different sequences a encoding different protein, therefore the resulting immune response will also be different.

The special technical feature of group 6, 18 and 33 is considered to be a method of producing a cellular mediated immune response to the heterologous protein encoded by the different adenoviral vectors in conjunction with immunizing the individual a DNA plasmid vaccine. Each method contains different sequences encoding a different protein, therefore the resulting immune response will also be different.

Accordingly, groups 1-48 are not so linked by the same or corresponding technical feature as to form a single general inventive concept.

Continuation of B. FIELDS SEARCHED Item 3:

WEST 2.0, STN-BIOSIS, MEDLINE

adenoviral vector, deletion, HIV, Gag, polyadenylation signal, CMV promoter

CORRECTED VERSION

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- (84) Designated States (regional): ARIPO patent (GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG).

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(54) Title: ENHANCED FIRST GENERATION ADENOVIRUS VACCINES EXPRESSING CODON OPTIMIZED HIV1-GAG, POL, NEF AND MODIFICATIONS

(57) Abstract: First generation adenoviral vectors and associated recombinant adenovirus-based HIV vaccines which show enhanced stability and growth properties and greater cellular-mediated immunity are described within this specification. These adenoviral vectors are utilized to generate and produce through cell culture various adenoviral-based HIV-1 vaccines which contain HIV-1 gag, HIV-1 pol and/or HIV-1 nef polynucleotide pharmaceutical products, and biologically relevant modifications thereof. These adenovirus vaccines, when directly introduced into living vertebrate tissue, preferably a mammalian host such as a human or a non-human mammal of commercial or domestic veterinary importance, express the HIV1- Gag, Pol and/or Nef protein or biologically modification thereof, inducing a cellular immune response which specifically recognizes HIV-1. The exemplified polynucleotides of the present invention are synthetic DNA molecules encoding HIV-1 Gag, encoding codon optimized HIV-1 Pol, derivatives of optimized HIV-1 Pol (including constructs wherein protease, reverse transcriptase, RNAse H and integrase activity of HIV-1 Pol is inactivated), HIV-1 Nef and derivatives of optimized HIV-1 Nef, including nef mutants which effect wild type characteristics of Nef, such as myristylation and down regulation of host CD4. The adenoviral vaccines of the present invention, when administered alone or in a combined modality regime, will offer a prophylactic advantage to previously uninfected individuals and/or provide a therapeutic effect by reducing viral load levels within an infected individual, thus prolonging the asymptomatic phase of HIV-1 infection.



For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

TITLE OF THE INVENTION

ENHANCED FIRST GENERATION ADENOVIRUS VACCINES EXPRESSING CODON OPTIMIZED HIV1-GAG, POL, NEF AND MODIFICATIONS

5 CROSS-REFERENCE TO RELATED APPLICATIONS

This application claims the benefit, under 35 U.S.C. §119(e), of U.S. provisional applications 60/233,180, 60/279,056, and Attorney Docket 20867PV2 (serial number unassigned), filed September 15, 2000, March 27, 2001, and September 7, 2001, respectively.

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STATEMENT REGARDING FEDERALLY-SPONSORED R&D Not Applicable

REFERENCE TO MICROFICHE APPENDIX

Not Applicable

FIELD OF THE INVENTION

The present invention relates to recombinant, replication-deficient first generation adenovirus vaccines found to exhibit enhanced growth properties and greater cellular-mediated immunity as compared to other replication-deficient vectors. The invention also relates to the associated first generation adenoviral vectors described herein, which, through the incorporation of additional 5' adenovirus sequence, enhance large scale production efficiency of the recombinant, replicationdefective adenovirus described herein. Another aspect of the instant invention is the surprising discovery that the intron A portion of the human cytomegalovirus (hCMV) promoter constitutes a region of instability in adenoviral vector constructs. Removal of this region from adenoviral expression constructs results in greatly improved vector stability. Therefore, improved vectors expressing a transgene under the control of an intron A-deleted CMV promoter constitute a further aspect of this invention. These adenoviral vectors are useful for generating recombinant adenovirus vaccines against human immunodeficiency virus (HIV). In particular, the first generation adenovirus vectors disclosed herein are utilized to construct and generate adenovirus-based HIV-1 vaccines which contain HIV-1 Gag, HIV-1 Pol and/or HIV-1 Nef polynucleotide pharmaceutical products, and biologically active modifications thereof. Host administration of the recombinant, replication-deficient adenovirus vaccines described herein results in expression of HIV-1 Gag, HIV-1- Pol and/or Nef protein or

immunologically relevant modifications thereof, inducing a cellular immune response which specifically recognizes HIV-1. The exemplified polynucleotides of the present invention are synthetic DNA molecules encoding codon optimized HIV-1 Gag, HIV-1 Pol, derivatives of optimized HIV-1 Pol (including constructs wherein protease, reverse transcriptase, RNAse H and integrase activity of HIV-1 Pol is inactivated), HIV-1 Nef, and derivatives of optimized HIV-1 Nef, including nef mutants which effect wild type characteristics of Nef, such as myristylation and down regulation of host CD4. The HIV adenovirus vaccines of the present invention, when administered alone or in a combined modality and/or prime/boost regimen, will offer a prophylactic advantage to previously uninfected individuals and/or provide a therapeutic effect by reducing viral load levels within an infected individual, thus prolonging the asymptomatic phase of HIV-1 infection.

BACKGROUND OF THE INVENTION

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Human Immunodeficiency Virus-1 (HIV-1) is the etiological agent of acquired human immune deficiency syndrome (AIDS) and related disorders. HIV-1 is an RNA virus of the Retroviridae family and exhibits the 5'LTR-gag-pol-env-LTR 3'organization of all retroviruses. The integrated form of HIV-1, known as the provirus, is approximately 9.8 Kb in length. Each end of the viral genome contains flanking sequences known as long terminal repeats (LTRs). The HIV genes encode at least nine proteins and are divided into three classes; the major structural proteins (Gag, Pol, and Env), the regulatory proteins (Tat and Rev); and the accessory proteins (Vpu, Vpr, Vif and Nef).

The gag gene encodes a 55-kilodalton (kDa) precursor protein (p55) which is expressed from the unspliced viral mRNA and is proteolytically processed by the HIV protease, a product of the pol gene. The mature p55 protein products are p17 (matrix), p24 (capsid), p9 (nucleocapsid) and p6.

The pol gene encodes proteins necessary for virus replication; a reverse transcriptase, a protease, integrase and RNAse H. These viral proteins are expressed as a Gag-Pol fusion protein, a 160 kDa precursor protein which is generated via a ribosomal frame shifting. The viral encoded protease proteolytically cleaves the Pol polypeptide away from the Gag-Pol fusion and further cleaves the Pol polypeptide to the mature proteins which provide protease (Pro, P10), reverse transcriptase (RT, P50), integrase (IN, p31) and RNAse H (RNAse, p15) activities.

The *nef* gene encodes an early accessory HIV protein (Nef) which has been shown to possess several activities such as down regulating CD4 expression, disturbing T-cell activation and stimulating HIV infectivity.

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The env gene encodes the viral envelope glycoprotein that is translated as a 160-kilodalton (kDa) precursor (gp160) and then cleaved by a cellular protease to yield the external 120-kDa envelope glycoprotein (gp120) and the transmembrane 41-kDa envelope glycoprotein (gp41). Gp120 and gp41 remain associated and are displayed on the viral particles and the surface of HIV-infected cells.

The tat gene encodes a long form and a short form of the Tat protein, a RNA binding protein which is a transcriptional transactivator essential for HIV-1 replication.

The rev gene encodes the 13 kDa Rev protein, a RNA binding protein. The Rev protein binds to a region of the viral RNA termed the Rev response element (RRE). The Rev protein promotes transfer of unspliced viral RNA from the nucleus to the cytoplasm. The Rev protein is required for HIV late gene expression and in turn, HIV replication.

Gp120 binds to the CD4/chemokine receptor present on the surface of helper T-lymphocytes, macrophages and other target cells in addition to other co-receptor molecules. X4 (macrophage tropic) virus show tropism for CD4/CXCR4 complexes while a R5 (T-cell line tropic) virus interacts with a CD4/CCR5 receptor complex. After gp120 binds to CD4, gp41 mediates the fusion event responsible for virus entry. The virus fuses with and enters the target cell, followed by reverse transcription of its single stranded RNA genome into the double-stranded DNA via a RNA dependent DNA polymerase. The viral DNA, known as provirus, enters the cell nucleus, where the viral DNA directs the production of new viral RNA within the nucleus, expression of early and late HIV viral proteins, and subsequently the production and cellular release of new virus particles. Recent advances in the ability to detect viral load within the host shows that the primary infection results in an extremely high generation and tissue distribution of the virus, followed by a steady state level of virus (albeit through a continual viral production and turnover during this phase), leading ultimately to another burst of virus load which leads to the onset of clinical AIDS. Productively infected cells have a half life of several days, whereas chronically or latently infected cells have a 3-week half life, followed by non-productively infected cells which have a long half life (over 100 days) but do not significantly contribute to day to day viral loads seen throughout the course of disease.

Destruction of CD4 helper T lymphocytes, which are critical to immune defense, is a major cause of the progressive immune dysfunction that is the hallmark of HIV infection. The loss of CD4 T-cells seriously impairs the body's ability to fight most invaders, but it has a particularly severe impact on the defenses against viruses, fungi, parasites and certain bacteria, including mycobacteria.

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Effective treatment regimens for HIV-1 infected individuals have become available recently. However, these drugs will not have a significant impact on the disease in many parts of the world and they will have a minimal impact in halting the spread of infection within the human population. As is true of many other infectious diseases, a significant epidemiologic impact on the spread of HIV-1 infection will only occur subsequent to the development and introduction of an effective vaccine. There are a number of factors that have contributed to the lack of successful vaccine development to date. As noted above, it is now apparent that in a chronically infected person there exists constant virus production in spite of the presence of anti-HIV-1 humoral and cellular immune responses and destruction of virally infected cells. As in the case of other infectious diseases, the outcome of disease is the result of a balance between the kinetics and the magnitude of the immune response and the pathogen replicative rate and accessibility to the immune response. Pre-existing immunity may be more successful with an acute infection than an evolving immune response can be with an established infection. A second factor is the considerable genetic variability of the virus. Although anti-HTV-1 antibodies exist that can neutralize HIV-1 infectivity in cell culture, these antibodies are generally virus isolate-specific in their activity. It has proven impossible to define serological groupings of HIV-1 using traditional methods. Rather, the virus seems to define a serological "continuum" so that individual neutralizing antibody responses, at best, are effective against only a handful of viral variants. Given this latter observation, it would be useful to identify immunogens and related delivery technologies that are likely to elicit anti-HIV-1 cellular immune responses. It is known that in order to generate CTL responses antigen must be synthesized within or introduced into cells, subsequently processed into small peptides by the proteasome complex, and translocated into the endoplasmic reticulum/Golgi complex secretory pathway for eventual association with major histocompatibility complex (MHC) class I proteins. CD8⁺ T lymphocytes recognize antigen in association with class I MHC via the T cell receptor (TCR) and the CD8 cell surface protein. Activation of naive CD8⁺ T cells into activated effector or memory cells generally requires both TCR engagement of antigen as described above as well as engagement of costimulatory proteins. Optimal

induction of CTL responses usually requires "help" in the form of cytokines from CD4⁺ T lymphocytes which recognize antigen associated with MHC class II molecules via TCR and CD4 engagement.

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European Patent Applications 0 638 316 (Published February 15, 1995) and 0 586 076 (Published March 9, 1994), (both assigned to American Home Products Corporation) describe replicating adenovirus vectors carrying an HIV gene, including env or gag. Various treatment regimens were used with chimpanzees and dogs, some of which included booster adenovirus or protein plus alum treatments.

Replication-defective adenoviral vectors harboring deletions in the E1 region are known, and recent adenoviral vectors have incorporated the known packaging repeats into these vectors; e.g., see EP 0 707 071, disclosing, *inter alia*, an adenoviral vector deleted of E1 sequences from base pairs 459 to 3328; and U.S. Patent No. 6,033,908, disclosing, *inter alia*, an adenoviral vector deleted of base pairs 459-3510. The packaging efficiency of adenovirus has been taught to depend on the number of incorporated individual A (packaging) repeats; *see*, *e.g.*, Gräble and Hearing, 1990 *J. Virol.* 64(5):2047-2056; Gräble and Hearing, 1992 *J. Virol.* 66(2):723-731.

Larder, et al., (1987, Nature 327: 716-717) and Larder, et al., (1989, Proc. Natl. Acad. Sci. 86: 4803-4807) disclose site specific mutagenesis of HIV-1 RT and the effect such changes have on *in vitro* activity and infectivity related to interaction with known inhibitors of RT.

Davies, et al. (1991, *Science* 252:, 88-95) disclose the crystal structure of the RNase H domain of HIV-1 Pol.

Schatz, et al. (1989, FEBS Lett. 257: 311-314) disclose that mutations Glu478Gln and His539Phe in a complete HIV-1 RT/RNase H DNA fragment results in defective RNase activity without effecting RT activity.

Mizrahi, et al. (1990, *Nucl. Acids. Res.* 18: pp. 5359-5353) disclose additional mutations Asp443Asn and Asp498Asn in the RNase region of the *pol* gene which also results in defective RNase activity. The authors note that the Asp498Asn mutant was difficult to characterize due to instability of this mutant protein.

Leavitt, et al. (1993, J. Biol. Chem. 268: 2113-2119) disclose several mutations, including a Asp64Val mutation, which show differing effect on HIV-1 integrase (IN) activity.

Wiskerchen, et al. (1995, J. Virol. 69: 376-386) disclose singe and double mutants, including mutation of aspartic acid residues which effect HIV-1 IN and viral replication functions.

It would be of great import in the battle against AIDS to produce a prophylactic- and/or therapeutic-based HIV vaccine which generates a strong cellular immune response against an HIV infection. The present invention addresses and meets these needs by disclosing a class of adenovirus vaccines which, upon host administration, express codon optimized and modified versions of the HIV-1 genes, gag, pol and nef. These recombinant, replication-defective adenovirus vaccines may be administered to a host, such as a human, alone or as part of a combined modality regimen and/or prime-boost vaccination regimen with components of the present invention and/or a distinct viral HIV DNA vaccine, non-viral HIV DNA vaccine, HIV subunit vaccine, an HIV whole killed vaccine and/or a live attenuated HIV vaccine.

SUMMARY OF THE INVENTION

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The present invention relates to enhanced replication-defective recombinant adenovirus vaccine vectors and associated recombinant, replication-deficient adenovirus vaccines which encode various forms of HIV-1 Gag, HIV-1 Pol, and/or HIV-1 Nef, including immunologically relevant modifications of HIV-1 Gag, HIV-1 Pol and HIV-1 Nef. The adenovirus vaccines of the present invention express HIV antigens and provide for improved cellular-mediated immune responses upon host administration. Potential vaccinees include but are not limited to primates and especially humans and non-human primates, and also include any non-human mammal of commercial or domestic veterinary importance. An effect of the improved recombinant adenovirus-based vaccines of the present invention should be a lower transmission rate to previously uninfected individuals (i.e., prophylactic applications) and/or reduction in the levels of the viral loads within an infected individual (i.e., therapeutic applications), so as to prolong the asymptomatic phase of HIV-1 infection. In particular, the present invention relates to adenoviral-based vaccines which encode various forms of codon optimized HIV-1 Gag (including but in no way limited to p55 versions of codon optimized full length (FL) Gag and tPA-Gag fusion proteins), HIV-1 Pol, HIV-1 Nef, and selected modifications of immunological relevance. The administration, intracellular delivery and expression of these adenovirus vaccines elicit a host CTL and Th response. The preferred replication-defective recombinant adenoviral vaccine vectors include but are not limited to synthetic DNA molecules which (1) encode codon optimized versions of wild type HIV-1 Gag; (2) encode codon optimized versions of HIV-1 Pol; (3) encode codon optimized versions of HIV-1 Pol fusion proteins; (4) encode codon optimized versions of modified HIV-1 Pol proteins and fusion proteins, including but not limited

to pol modifications involving residues within the catalytic regions responsible for RT, RNase and IN activity within the host cell; (5) encode codon optimized versions of wild type HIV-1 Nef; (6) codon optimized versions of HIV-1 Nef fusion proteins; and/or (7) codon optimized versions of HIV-1 Nef derivatives, including but not limited to nef modifications involving introduction of an amino-terminal leader sequence, removal of an amino-terminal myristylation site and/or introduction of dileucine motif mutations. The Nef-based fusion and modified proteins, disclosed within this specification and expressed from an adenoviral-based vector vaccine this specification, may possess altered trafficking and/or host cell function while retaining the ability to be properly presented to the host MHC I complex and in turn elicit a host CTL and Th response. Examples of HIV-1 Gag, Pol and/or Nef fusion proteins include but are not limited to fusion of a leader or signal peptide at the NH₂-teriminal portion of the viral antigen coding region. Such a leader peptide includes but is not limited to a tPA leader peptide.

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The adenoviral vector utilized in construction of the HIV-1 Gag-, HIV-1 Poland/or HIV-1 Nef- based vaccines of the present invention may comprise any replication-defective adenoviral vector which provides for enhanced genetic stability of the recombinant adenoviral genome through large scale production and purification of the recombinant virus. In other words, an HIV-1 Gag-, Pol- or Nef-based adenovirus vaccine of the present invention is a purified recombinant, replicationdefective adenovirus which is shown to be genetically stable through multiple passages in cell culture and remains so during large scale production and purification procedures. Such a recombinant adenovirus vector and harvested adenovirus vaccine lends itself to large scale dose filling and subsequent worldwide distribution procedures which will be demanded of an efficacious monovalent or multivalent HIV vaccine. The present invention meets this basic requirement with description of a replication-defective adenoviral vector and vectors derived therefrom, at least partially deleted in E1, comprising a wildtype adenovirus cis-acting packaging region from about base pair 1 to between from about base pair 342 (more preferably, 400) to about base pair 458 of the wildtype adenovirus genome. A preferred embodiment of the instant invention comprises base pairs 1-450 of a wildtype adenovirus. In other preferred embodiments, the replication -defective adenoviral vector has, in addition thereto, a region 3' to the E1-deleted region comprising base pairs 3511-3523. Basepairs 342-450 (more particularly, 400-450) constitute an extension of the 5'region of previously disclosed vectors carrying viral antigens, particularly HIV antigens (see, e.g., PCT International Application PCT/US00/18332, published

January 11, 2001 (WO 01/02067), which claims priority to U.S. Provisional Application Serial Nos. 60/142,631 and 60/148,981, filed 7/6/1999 and 8/13/1999, respectively; these documents herein incorporated by reference. Applicants have found that extending the 5' region further into the E1 gene into the disclosed vaccine vectors incorporated elements found to be important in optimizing the packaging of the virus.

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As compared to previous vectors not comprising basepairs from about 1 to between from about base pair 342 (more preferably, 400) to about base pair 458 of the wildtype adenovirus genome, vectors comprising the above region exhibited enhanced 10 growth characteristics, with approximately 5-10 fold greater amplification rates, a more potent virus effect, allowing lower doses of virus to be used to generate equivalent immunity; and a greater cellular-mediated immune response than replication-deficient vectors not comprising this region (basepairs 1-450). Even more important, adenoviral constructs derived therefrom are very stable genetically in 15 large-scale production, particularly those comprising an expression cassette under the control of a hCMV promoter devoid of intron A. This is because Applicants have surprisingly found that the intron A portion of the hCMV promoter constituted a region of instability when employed in adenoviral vectors. Applicants have, therefore, identified an enhanced adenoviral vector which is particularly suited for use 20 in gene therapy and nucleotide-based vaccine-vectors which, favorably, lends itself to large scale propagation.

A preferred embodiment of this invention is a replication-defective adenoviral vector in accordance with the above description wherein the gene is inserted in the form of a gene expression cassette comprising (a) a nucleic acid encoding a protein or biologically active and/or immunologically relevant portion thereof; (b) a heterologous promoter operatively linked to the nucleic acid of part a); and, (c) a transcription terminator.

In preferred embodiments, the E1 gene, other than that contained within basepairs 1-450 or, alternatively, that contained within base pairs 1-450 and 3511-3523 has been deleted from the adenoviral vector, and the gene expression cassette has replaced the deleted E1 gene. In other preferred embodiments, the replication defective adenovirus genome does not have a functional E3 gene, or the E3 gene has been deleted. Most preferably, the E3 region is present within the adenoviral genome. Further preferred embodiments are wherein the gene expression cassette is in an E1 anti-parallel (transcribed in a 3' to 5' direction relative to the vector backbone)

orientation or, more preferably, an E1 parallel (transcribed in a 5' to 3' direction relative to the vector backbone) orientation.

Further embodiments relate to a shuttle plasmid vector comprising: an adenoviral portion and a plasmid portion, wherein said adenovirus portion comprises: a) a replication defective adenovirus genome, at least partially deleted in E1, comprising a wildtype adenovirus *cis*-acting packaging region from about base pair 1 to between from about base pair 342 (more preferably, 400) to about base pair 458 (preferably, 1-450) of the wildtype adenovirus genome and, preferably, in addition thereto, basepairs 3511-3523 of a wildtype adenovirus sequence; and b) a gene expression cassette comprising: (a) a nucleic acid encoding a protein or biologically active and/or immunologically relevant portion thereof; (b) a heterologous promoter operatively linked to the nucleic acid of part a); and (c) a transcription terminator and/or a polyadenylation site.

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Other aspects of this invention include a host cell comprising said adenoviral vectors and/or said shuttle plasmid vectors; vaccine compositions comprising said vectors; and methods of producing the vectors comprising (a) introducing the adenoviral vector into a host cell which expresses adenoviral E1 protein, and (b) harvesting the resultant adenoviral vectors.

To this end, the present invention particularly relates to harvested recombinant, replication defective virus derived from a host cell, such as but not limited to 293 cells or PER.C6® cells, including but not limited to harvested virus related to any of the MRKAd5 vector backbones, with or without an accompanying transgene, including but not limited to the HIV-1 antigens described herein. An HIV-1 vaccine is represented by any harvested, recombinant adenovirus material which expresses any one or more of the HIV-1 antigens disclosed herein. This harvested material may then be purified, formulated and stored prior to host administration.

Another aspect of this invention is a method of generating a cellular immune response against a protein in an individual comprising administering to the individual an adenovirus vaccine vector comprising:

a) a recombinant, replication defective adenoviral vector, at least partially deleted in E1, comprising a wildtype adenovirus *cis*-acting adenovirus packaging region from about base pair 1 to between from about base pair 342 (more preferably, 400) to about base pair 458 (preferably, 1-450) and, preferably in addition thereto, base pairs 3511-3523 of a wildtype adenovirus sequence, and,

b) a gene expression cassette comprising:(i) a nucleic acid encoding a protein or biologically active and/or immunologically relevant portion thereof; (ii) a heterologous promoter operatively linked to the nucleic acid of part a); and (iii) a transcription terminator and/or a polyadenylation site.

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In view of the efficacious nature of the adenoviral and/or DNA plasmid vaccines described herein, the present invention relates to all methodology regarding administration of one or more of these adenoviral and/or DNA plasmid vaccines to provide effective immunoprophylaxis, to prevent establishment of an HIV-1 infection following exposure to this virus, or as a post-HIV infection therapeutic vaccine to mitigate the acute HIV-1 infection so as to result in the establishment of a lower virus load with beneficial long term consequences. As discussed herein, such a treatment regimen may include a monovalent or multivalent composition, various combined modality applications, and/or a prime/boost regimen to as to optimize antigen expression and a concomitant cellular-mediated and/or humoral immune response upon inoculation into a living vertebrate tissue. Therefore, the present invention provides for methods of using the adenoviral and/or DNA plasmid vaccines disclosed herein within the various parameters disclosed herein as well as any additional parameters known in the art, which, upon introduction into mammalian tissue induces intracellular expression of the gag, pol and/or nef-based vaccines.

To this end, the present invention relates in part to methods of generating a cellular immune response in a vaccinee, preferably a human vaccinee, wherein the individual is given more than one administration of adenovirus vaccine vector, and it may be given in a regimen accompanied by the administration of a plasmid vaccine. The plasmid vaccine (also referred to herein as a "DNA plasmid vaccine" or "vaccine plasmid" comprises a nucleic acid encoding a protein or an immunologically relevant portion thereof, a heterologous promoter operably linked to the nucleic acid sequence, and a transcription terminator or a polyadenylation signal (such as bGH or SPA, respectively). There may be a predetermined minimum amount of time separating the administrations. The individual can be given a first dose of plasmid vaccine, and then a second dose of plasmid vaccine. Alternatively, the individual may be given a first dose of adenovirus vaccine, and then a second dose of adenovirus vaccine. In other embodiments, the plasmid vaccine is administered first, followed after a time by administration of the adenovirus vaccine. Conversely, the adenovirus vaccine may be administered first, followed by administration of plasmid vaccine after a time. In these embodiments, an individual may be given multiple doses of the same adenovirus serotype in either viral vector or plasmid form, or the virus may be of

differing serotypes. In the alternative, a viral antigen of interest can be first delivered via a viral vaccine other than an adenovirus-based vaccine, and then followed with the adenoviral vaccine disclosed. Alternative viral vaccines include but are not limited to pox virus and venezuelan equine encephilitis virus.

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The present invention also relates to multivalent adenovirus vaccine compositions which comprise Gag, Pol and Nef components described herein; see, e.g., Example 29 and Table 25. Such compositions will provide for an enhanced cellular immune response subsequent to host administration, particularly given the genetic diversity of human MHCs and of circulating virus. Examples, but not limitations, include MRKAd5-vector based multivalent vaccine compositions which provide for a divalent (i.e., gag and nef, gag and pol, or pol and nef components) or a trivalent vaccine (i.e., gag, pol and nef components) composition. Such a mutlivalent vaccine may be filled for a single dose or may consist of multiple inoculations of each individually filled component; and may in addition be part of a prime/boost regimen with viral or non-viral vector vaccines as introduced in the previous paragraph. To this end, preferred compositions are MRKAd5 adenovirus used in combination with multiple, distinct HIV antigen classes. Each HIV antigen class is subject to sequence manipulation, thus providing for a multitude of potential vaccine combinations; and such combinations are within the scope of the present invention. The utilization of such combined modalities vaccine formulation and administration increase the probability of eliciting an even more potent cellular immune response when compared to inoculation with a single modality regimen.

The concept of a "combined modality" as disclosed herein also covers the alternative mode of administration whereby multiple HIV-1 viral antigens may be ligated into a proper shuttle plasmid for generation of a pre-adenoviral plasmid comprising multiple open reading frames. For example, a trivalent vector may comprise a gag-pol-nef fusion, in either a E3(-) or E3(+) background, preferably a E3 deleted backbone, or possibly a "2+1" divalent vaccine, such as a gag-pol fusion (i.e., codon optimized p55 gag and inactivated optimized pol; Example 29 and Table 25) within the same MRKAd5 backbone, with each open reading frame being operatively linked to a distinct promoter and transcription termination sequence. Alternatively, the two open reading frames may be operatively linked to a single promoter, with the open reading frames operatively linked by an internal ribosome entry sequence (IRES). Therefore, a multivalent vaccine delivered as a single, or possibly a second harvested recombinant, replication-deficient adenovirus is contemplated as part of the present invention.

Therefore, the adenoviral vaccines and plasmid DNA vaccines of this invention may be administered alone, or may be part of a prime and boost administration regimen. A mixed modality priming and booster inoculation scheme will result in an enhanced immune response, particularly if pre-existing anti-vector immune responses are present. This one aspect of this invention is a method of priming a subject with the plasmid vaccine by administering the plasmid vaccine at least one time, allowing a predetermined length of time to pass, and then boosting by administering the adenoviral vaccine. Multiple primings typically, 1-4, are usually employed, although more may be used. The length of time between priming and boost may typically vary from about four months to a year, but other time frames may be used. In experiments with rhesus monkeys, the animals were primed four times with plasmid vaccines, then were boosted 4 months later with the adenoviral vaccine. Their cellular immune response was notably higher than that of animals which had only received adenoviral vaccine. The use of a priming regimen may be particularly preferred in situations where a person has a pre-existing anti-adenovirus immune response.

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It is an object of the present invention to provide for enhanced replication-defective recombinant adenoviral vaccine vector backbones. These recombinant adenoviral backbones may accept one or more transgenes, which may be passaged through cell culture for growth, amplification and harvest.

It is a further object to provide for enhanced replication-defective recombinant adenoviral vaccine vectors which encode various transgenes.

It is also an object of the present invention to provide for a harvested recombinant, replication-deficient adenovirus which shows enhanced growth and amplification rates while in combination with increased virus stability after continuous passage in cell culture. Such a recombinant adenovirus is particularly suited for use in gene therapy and nucleotide-based vaccine vectors which, favorably, lends itself to large scale propagation.

To this end, it is an object of the present invention to provide for (1) enhanced replication-defective recombinant adenoviral vaccine vectors as described herein which encode various forms of HIV-1 Gag, HIV-1 Pol, and/or HIV-1 Nef, including immunologically relevant modifications of HIV-1 Gag, HIV-1 Pol and HIV-1 Nef, and (2) harvested, purified recombinant replication-deficient adenovirus generated by passage of the adenoviral vectors of (1) through one or multiple passages through cell culture, including but not limited to passage through 293 cells or PER.C6® cells.

It is also an object of the present invention to provide for recombinant adenovirus harvested by one or multiple passages through cell culture. As relating to recombinant adenoviral vaccine vector, this recombinant virus is harvested and formulated for subsequent host administration.

It is also an object of the present invention to provide for replication-defective adenoviral vectors wherein at least one gene is inserted in the form of a gene expression cassette comprising (a) a nucleic acid encoding a protein or biologically active and/or immunologically relevant portion thereof; (b) a heterologous promoter operatively linked to the nucleic acid of part a); and, (c) a transcription terminator.

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It is also an object of the present invention to provide for a host cell comprising said adenoviral vectors and/or said shuttle plasmid vectors; vaccine compositions comprising said vectors; and methods of producing the vectors comprising (a) introducing the adenoviral vector into a host cell which expresses adenoviral E1 protein, and (b) harvesting the resultant adenoviral vectors. It is a further object of the present invention to provide for methods of generating a cellular immune response against a protein in an individual comprising administering to the individual an adenovirus vaccine vector comprising a) a replication defective adenoviral vector, at least partially deleted in E1, comprising a wildtype adenovirus cis-acting packaging region from about base pair 1 to between from about base pair 342 (more preferably, 400) to about 450 (preferably, 1-450) and, preferably, 3511-3523 of a wildtype adenovirus sequence, and, b) a gene expression cassette comprising:(i) a nucleic acid encoding a protein or biologically active and/or immunologically relevant portion thereof; (ii) a heterologous promoter operatively linked to the nucleic acid of part a); and (iii) a transcription terminator and/or a polyadenylation site.

It is also an object of the present invention to provide various alternatives for vaccine administration regimes, namely administration of one or more adenoviral and/or DNA plasmid vaccines described herein to provide effective immunoprophylaxis for uninfected individuals or a therapeutic treatment for HIV infected patients. Such processes include but are not limited to multivalent HIV-1 vaccine compositions, various combined modality regimes as well as various prime/boost alternatives. These methods of administration, relating to vaccine composition and/or scheduled administration, will increase the probability of eliciting an even more potent cellular immune response when compared to inoculation with a single modality regimen.

As used throughout the specification and claims, the following definitions and abbreviations are used:

"HAART" refers to - highly active antiretroviral therapy -.

"first generation" vectors are characterized as being replication-defective.

They typically have a deleted or inactivated E1 gene region, and preferably have a deleted or inactivated E3 gene region as well.

"AEX" refers to Anion Exchange chromatography.

"QPA" refers to Quick PCR-based Potency Assay.

"bps" refers to basepairs.

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"s" or "str" denotes that the transgene is in the E1 parallel or "straight" orientation.

"PBMCs" refers to peripheral blood monocyte cells.

"FL" refers to full length.

"FLgag" refers to a full-length optimized gag gene, as shown in Figure 2.

"Ad5-Flgag" refers to an adenovirus serotype 5 replication deficient virus which carries an expression cassette which comprises a full length optimized gag gene under the control of a CMV promoter.

"Promoter" means a recognition site on a DNA strand to which an RNA polymerase binds. The promoter forms an initiation complex with RNA polymerase to initiate and drive transcriptional activity. The complex can be modified by activating sequences such as enhancers or inhibiting sequences such as silencers.

"Leader" means a DNA sequence at the 5' end of a structural gene which is transcribed along with the gene. This usually results a protein having an N-terminal peptide extension, often referred to as a pro-sequences.

"Intron" means a section of DNA occurring in the middle of a gene which does not code for an amino acid in the gene product. The precursor RNA of the intron is excised and is therefore not transcribed into mRNA not translated into protein.

"Immunologically relevant" or "biologically active" means (1) with regards to a viral protein, that the protein is capable, upon administration, of eliciting a measurable immune response within an individual sufficient to retard the propagation and/or spread of the virus and/or to reduce the viral load present within the individual; or (2) with regards to a nucleotide sequence, that the sequence is capable of encoding for a protein capable of the above.

"Cassette" refers to a nucleic acid sequence which is to be expressed, along with its transcription and translational control sequences. By changing the cassette, a vector can express a different sequence.

"bGHpA" refers to the bovine growth hormone transcription terminator/polyadenylation sequence.

"tPAgag" refers to a fusion between the leader sequence of the tissue plasminogen activator leader sequence and an optimized HIV gag gene, as exemplified in Figure 30A-B, whether in a DNA or adenovirus-based vaccine vector.

Where utilized, "IA" or "inact" refers to an <u>inactivated</u> version of a gene (e.g. IApol).

"MCS" is "multiple cloning site".

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In general, adenoviral constructs, gene constructs are named by reference to the genes contained therein. For example:

"Ad5 HIV-1 gag", also referred to as the original HIV-1 gag adenoviral vector, is a vector containing a transgene cassette composed of a hCMV intron A promoter, the full length version of the human codon-optimized HIV-1 gag gene, and the bovine growth hormone polyadenylation signal. The transgene was inserted in the E1 antiparallel orientation in an E1 and E3 deleted adenovector.

"MRK Ad5 HIV-1 gag" also referred to as "MRKAd5gag" or "Ad5gag2" is an adenoviral vector taught herein which is deleted of E1, comprises basepairs 1-450 and 3511-3523, and has a human codon-optimized HIV-1 gene in an E1 parallel orientation under the control of a CMV promoter without intron A. The construct also comprises a bovine growth hormone polyadenylation signal.

"pV1JnsHIVgag", also referred to as "HIVFLgagPR9901", is a plasmid comprising the CMV immediate-early (IE) promoter and intron A, a full-length codon-optimized HIV gag gene, a bovine growth hormone-derived polyadenylation and transcriptional termination sequence, and a minimal pUC backbone.

"pV1JnsCMV(no intron)-FLgag-bGHpA" is a plasmid derived from pV1JnsHIVgag which is deleted of the intron A portion of CMV and which comprises the full length HIV gag gene. This plasmid is also referred to as "pV1JnsHIVgag-bGHpA", pV1Jns-hCMV-FL-gag-bGHpA" and "pV1JnsCMV(no intron) + FLgag + bGHpA".

"pV1JnsCMV(no intron)-FLgag-SPA" is a plasmid of the same composition as pV1JnsCMV(no intron)-FLgag-bGHpA except that the SPA termination sequence replaces that of bGHpA. This plasmid is also referred to as "pV1Jns-HIVgag-SPA" and pV1Jns-hCMV-FLgag-SPA".

"pdelE1sp1A" is a universal shuttle vector with no expression cassette (i.e., no promoter or polyA). The vector comprises wildtype adenovirus serotype 5 (Ad5) sequences from bp 1 to bp 341 and bp 3524 to bp 5798, and has a multiple cloning

site between the Ad5 sequences ending 341 bp and beginning 3524 bp. This plasmid is also referred to as the original Ad 5 shuttle vector.

"MRKpdelE1sp1A" or "MRKpdelE1(Pac/pIX/pack450)" or

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"MRKpdelE1(Pac/pIX/pack450)Cla1" is a universal shuttle vector with no expression cassette (i.e. no promoter or polyA) comprising wildtype adenovirus serotype 5 (Ad5) sequences from bp1 to bp450 and bp 3511 to bp 5798. The vector has a multiple cloning site between the Ad5 sequence ending 450 bp and beginning 3511 bp. This shuttle vector may be used to insert the CMV promoter and the bGHpA fragments in both the straight ("str". or E1 parallel) orientation or in the opposite (opp. or E1 antiparallel) orientation)

"MRKpdelE1(Pac/pIX/pack450)+CMVmin+BGHpA(str.)" is still another shuttle vector which is the modified vector that contains the CMV promoter (no intronA) and the bGHpA fragments. The expression unit containing the hCMV promoter (no intron A) and the bovine growth hormone polyadenylation signal has been inserted into the shuttle vector such that insertion of the gene of choice at a unique BgIII site will ensure the direction of transcription of the transgene will be Ad5 E1 parallel when inserted into the MRKpAd5(E1/E3+)Cla1 pre-plasmid. This shuttle vector, as shown in Figures 22 and 23, was used to insert the respective IApol and G2A,LLAA nef genes directly into.

"MRKpdelE1-CMV(no intron)-FLgag-bGHpA" is a shuttle comprising Ad5 sequences from basepairs 1-450 and 3511-5798, with an expression cassette containing human CMV without intron A, the full-length human codon-optimized HIV gag gene and bovine growth hormone polyadenylation signal. This plasmid is also referred to as "MRKpdelE1 shuttle +hCMV-FL-gag-BGHpA"

"MRKpAdHVE3+CMV(no intron)-FLgag-bGHpA" is an adenoviral vector comprising all Ad5 sequences except those nucleotides encompassing the E1 region (from 451-3510), a human CMV promoter without intron A, a full-length human codon-optimized HIV gag gene, and a bovine growth hormone polyadenylation signal. This vector is also referred to as "MRKpAdHVE3 + hCMV-FL-gag-BGHpA", "MRKpAd5HIV-1gag", "MRKpAd5gag", "pMRKAd5gag" or "pAd5gag2".

"pV1Jns-HIV-pol inact(opt)" or "pV1Jns-HIV IA pol (opt) is the inactivated Pol gene (contained within SEQ ID NO:3) cloned into the BglII site of V1Jns (Figure 17A-C). As noted herein, various derivatives of HIV-1 pol may be cloned into a plasmid expression vector such as V1Jns or V1Jns-tPA, thus serving directly as DNA vaccine candidates or as a source for subcloning into an appropriate adenoviral vector.

"MRKpdel+hCMVmin+FL-pol+bGHpA(s)" is the "MRKpdelE1(Pac/pIX/pack450)+CMVmin+BGHpA(str.)" shuttle mentioned above which contains the IA pol gene is the proper orientation. This shuttle vector is used in a bacterial recombination with MRKpAd(E1-/E3+)Cla1.

"MRKpAd+hCMVmin+FL-pol+bGHpA(S)E3+", also referred to herein as "pMRKAd5pol", is the pre-adenovirus plasmid which comprises a CMV-pol inact(opt)-pGHpA construct. The construction of this pre-adenovirus plasmid is shown in Figure 22.

"pV1Jns/nef (G2A,LLAA)" or "V1Jns/opt nef (G2A,LLAA)" comprises codon optimized HIV-1 Nef wherein the open reading frame codes for modifications at the amino terminal myristylation site (Gly-2 to Ala-2) and substitution of the Leu-174-Leu-175 dileucine motif to Ala-174-Ala-175 (SEQ ID NO:13; which comprises an initiating methionine residue at nucleotides 12-14 and a "TAA" stop codon from nucleotides 660-662). This fragment is subcloned into the Bgl II site of V1Jns and/orV1Jns-tPA (Figures 16A-B). As noted above for HIV-1 pol, HIV-1 nef constructs may be cloned into a plasmid expression vector such as V1Jns or V1Jns-tPA, thus serving directly as DNA vaccine candidates or as a source for subcloning into an appropriate adenoviral vector.

"MRKpdelE1hCMVminFL-nefBGHpA(s)", also referred to herein as "pMRKAd5nef", is the pre-adenovirus plasmid which comprises a CMV-nef (G2A,LLAA) codon optimized sequence. The construction of this pre-adenovirus plasmid is shown in Figure 23.

BRIEF DESCRIPTION OF THE FIGURES

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Figure 1 shows the original HIV-1 gag adenovector (Ad5HIV-1gag). This vector is disclosed in PCT International Application No. PCT/US00/18332 (WO 01/02607) filed July 3, 2000, claiming priority to U.S. Provisional Application Serial No. 60/142,631, filed July 6, 1999 and U.S. Application Serial No. 60/148,981, filed August 13, 1999, all three applications which are hereby incorporated by reference.

Figure 2 shows the nucleic acid sequence (SEQ ID NO: 29) of the optimized human HIV-1 gag open reading frame.

Figure 3 shows diagrammatically the new transgene constructs in comparison with the original gag transgene.

Figure 4 shows the modifications made to the original adenovector backbone in the generation of the novel vectors of the instant invention.

Figure 5 shows the virus mixing experiments that were carried out to determine the effects of the addition made to the packaging signal region (Expt. #1) and the E3 gene on viral growth (Expt. #2). The bars denote the region of modifications made to the E1 deletion.

Figure 6 shows an autoradiograph of viral DNA analysis following the viral mixing experiments described in Examples 6 and 7.

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Figures 7A, 7B and 7C are as follows: Figure 7A shows the hCMV-Flgag-bGHpA adenovectors constructed within the MRKpAdHVE3 and MRKpAdHVO adenovector backbones. Both E1 parallel and E1 antiparallel transgene orientation are represented. Figure 7B shows the hCMV-Flgag-SPA adenovectors constructed within the MRKpAdHVE3 and MRKpAdHVO adenovector backbones. Again, both E1 parallel and E1 antiparallel transgene orientation are represented. Figure 7C shows the mCMV-Flgag-bGHpA adenovectors constructed within the MRKpAdHVE3 and MRKpAdHVO adenovector backbones. Once again, both E1 parallel and E1 antiparallel transgene orientation are represented.

Figure 8A shows the experiment designed to test the effect of transgene orientation.

Figure 8B shows the experiments designed to test the effect of polyadenylation signal.

Figure 9 shows viral DNA from the four adenoviral vectors tested (Example 12) at P5, following *Bst*E11 digestion.

Figure 10 shows viral DNA analysis of passages 11 and 12 of MRKpAdHVE3, MRKAd5HIV-1gag, and MRKAd5HIV-1gagE3-.

Figure 11 shows viral DNA analysis (*Hind*III digestion) of passage 6 MRKpAdHVE3 and MRKAd5HIV-1gag used to initiate the viral competition study. The last two lanes are passage 11 analysis of duplicate passages of the competition study (each virus at MOI of 280 viral particles).

Figure 12 shows viral DNA analysis by *Hind* III digestion on high passage numbers for MRKAd5HIV-1gag in serum-containing media with collections made at specified times. The first lane shows the 1kb DNA size marker. The other lanes represent pre-plasmid control (digested with Pac1 and *Hind*III), MRKAd5HIV-1gag at P16, P19, and P21.

Figure 13 shows serum anti-p24 levels at 3 wks post i.m. immunization of balb/c mice (n=10) with varying doses of several Adgag constructs: (A) MRK Ad5 HIV-1 gag (through passage 5); (B) MRKAd5 hCMV-FLgag-bGHpA (E3-); (C) MRKAd5 hCMV-FLgag-SPA (E3+); (D) MRKAd5 mCMV-FLgag-bGHpA (E3+);

(E) research lot (293 cell-derived) of Ad5HIV-1 gag; and (F) clinical lot (Ad5gagFN0001) of Ad5HIV-1 gag. Reported are the geometric mean titers (GMT) for each cohort along with the standard error bars.

Figure 14 shows a restriction map of the pMRKAd5HIV-1gag vector.

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Figures 15A-X illustrates the nucleotide sequence of the pMRKAd5HIV-1gag vector (SEQ ID NO:27.[coding] and SEQ ID NO:28 [non-coding]).

Figures 16A-B shows a schematic representation of DNA vaccine expression vectors V1Jns (A) and V1Jns-tPA (B), which are utilized for HIV-1 gag, pol and nef constructs in various DNA/viral vector combined modality regimens as disclosed herein.

Figures 17A-C shows the nucleotide (SEQ ID NO:3) and amino acid sequence (SEQ ID NO:4) of IA-Pol. Underlined codons and amino acids denote mutations, as listed in Table 1.

Figure 18 shows codon optimized nucleotide and amino acid sequences through the fusion junction of tPA-pol inact(opt) (contained within SEQ ID NOs: 7 and 8, respectively). The underlined portion represents the NH₂-terminal region of IA-Pol.

Figures 19A-B show a nucleotide sequence comparison between wild type nef(jrfl) and codon optimized nef. The wild type nef gene from the jrfl isolate consists of 648 nucleotides capable of encoding a 216 amino acid polypeptide. WT, wild type sequence (SEQ ID NO:19); opt, codon-optimized sequence (contained within SEQ ID NO:1). The Nef amino acid sequence is shown in one-letter code (SEQ ID NO:2).

Figures 20A-C show nucleotide sequences at junctions between nef coding sequence and plasmid backbone of nef expression vectors V1Jns/nef (Figure 20A), V1Jns/nef(G2A,LLAA) (Figure 20B), V1Jns/tpanef (Figure 20C) and V1Jns/tpanef(LLAA) (Figure 20C, also). 5' and 3' flanking sequences of codon optimized nef or codon optimized nef mutant genes are indicated by bold/italic letters; nef and nef mutant coding sequences are indicated by plain letters. Also indicated (as underlined) are the restriction endonuclease sites involved in construction of respective nef expression vectors. V1Jns/tpanef and V1Jns/tpanef(LLAA) have identical sequences at the junctions.

Figure 21 shows a schematic presentation of nef and nef derivatives. Amino acid residues involved in Nef derivatives are presented. Glycine 2 and Leucine 174 and 175 are the sites involved in myristylation and dileucine motif, respectively. For both versions of the tpanef fusion genes, the putative leader peptide cleavage sites are

indicated with "*", and a exogenous serine residue introduced during the construction of the mutants is underlined.

Figure 22 shows diagrammatically the construction of the pre-adenovirus plasmid construct, MRKAd5Pol.

Figure 23 shows diagrammatically the construction of the pre-adenovirus plasmid construct, MRKAd5Nef.

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Figure 24 shows a comparison of clade B vs. clade C anti-gag T cell responses in clade B HIV-infected subjects.

Figure 25 shows a comparison of clade B vs. clade C anti-nef T cell responses in clade B HIV-infected subjects.

Figures 26A-AO illustrates the nucleotide sequence of the pMRKAd5HIV-1pol adenoviral vector (SEQ ID NO:32 [coding] and SEQ ID NO:33 [non-coding]), comprising the coding region of the inactivated pol gene (SEQ ID NO3).

Figures 27A-AM illustrates the nucleotide sequence of the pMRKAd5HIV-1 nef adenoviral vector (SEQ ID NO:34 [coding] and SEQ ID NO:35 [non-coding]), comprising the coding region of the inactivated pol gene (SEQ ID NO13).

Figure 28 shows the stability of MRKAd5 vectors comprising various promoter fragments (hCMV or mCMV) and terminations signals (bGH or SPA) in E3(+) or E3(-) backbones.

Figures 29A and B shows the anion-exchange HPLC viral particle concentrations of the freeze-thaw recovered cell associated virus at the 24, 36, 48, and 60 hpi time points (Figure 29A) and the timcourse QPA supernatant titers (Figure 29B) for MRKAd5gag, MRKAd5pol and MRKAd5nef.

Figure 30 shows the nucleotide sequence (SEQ ID NO:36) and amino acid sequence (SEQ ID NO:37) comprising the open reading frame of a representative tPA-gag fusion for use in the DNA and/or adenoviral vaccine disclosed herein.

Figure 31 shows the intracellular γIFN staining of PBMCs collected at week 10 (post DNA prime) and week 30 (post Ad boost). The cells were stimulated overnight in the presence or absence of the gag peptide pool. They were subsequently stained using fluorescence-tagged anti-CD3, anti-CD8, anti-CD4, and anti-γIFN monoclonal antibodies. Each plot shows all CD3+ T cells which were segregated in terms of positive staining for surface CD8 and γIFN production. The numbers in the upper right and lower right quadrants of each plot are the percentages of CD3⁺ cells that were CD8⁺γIFN⁺ and CD4⁺γIFN⁺, respectively.

Figure 32 shows a comparison of single-modality adenovirus immunization with DNA + adjuvant prime/adenovirus boost immunization.

Figures 33A-B show the nucleotide sequence (SEQ ID NO: 38) of the open reading frame for the gag-IApol fusion of Example 29.

Figures 34A-B show the protein sequence (SEQ ID NO:39) of the gag-IApol fustion frame.

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DETAILED DESCRIPTION OF THE INVENTION

A novel replication-defective, or "first generation," adenoviral vector suitable for use in gene therapy or nucleotide-based vaccine vectors is described. This vector is at least partially deleted in E1 and comprises a wildtype adenovirus cis-acting packaging region from about base pair 1 to between about base pair 342 (more preferably, 400) to about 458 (preferably, 1-450) and, preferably, 3511-3523 of a wild-type adenovirus sequence. It has been found that a vector of this description possesses enhanced growth characteristics, with approximately 5-10 fold greater amplification rates, and is more potent allowing lower doses of virus to be used to generate equivalent immunity. The vector, furthermore, generates a harvested recombinant adenovirus which shows greater cellular-mediated immune responses than replication-deficient vectors not comprising this region (basepairs 342-450). Adenoviral constructs derived from these vectors are, further, very stable genetically, particularly those comprising a transgene under the control of a hCMV promoter devoid of intron A. Viruses in accordance with this description were passaged continually and analyzed; see Example 12. Each virus analyzed maintained it correct genetic structure. Analysis was also carried out under propagation conditions similar to that performed in large scale production. Again, the vectors were found to possess enhanced genetic stability; see Figure 12. Following 21 passages, the viral DNA showed no evidence of rearrangement, and was highly reproducible from one production lot to the next. The outcome of all relevant tests indicate that the adenoviral vector is extremely well suited for large-scale production of recombinant, replication-deficient adenovirus, as shown herein with the data associated with Figure 28.

A preferred adenoviral vector in accordance with this description is a vector comprising basepairs 1-450, which is deleted in E3. This vector can accommodate up to approximately 7,500 base pairs of foreign DNA inserts (or exogenous genetic material). Another preferred vector is one retaining E3 which comprises basepairs 1-450. A preferred vector of this description is an E3+ vector comprising basepairs 1-450 and 3511-3523. This vector, when deleted of the region spanning basepairs 451-3510, can accommodate up to approximately, 4,850 base pairs of foreign DNA inserts

(or exogenous genetic material). The cloning capacities of the above vectors have been determined using 105% of the wildtype Ad5 sequence as the upper genome size limit.

Wildtype adenovirus serotype 5 is used as the basis for the specific basepair numbers provided throughout the specification. The wildtype adenovirus serotype 5 sequence is known and described in the art; see, Chroboczek et al., 1992 J. Virology 186:280, which is hereby incorporated by reference. Accordingly, a particular embodiment of the instant invention is a vector based on the adenovirus serotype 5 sequence. One of skill in the art can readily identify the above regions in other adenovirus serotypes (e.g., serotypes 2, 4, 6, 12, 16, 17, 24, 31, 33, and 42), regions defined by basepairs corresponding to the above basepair positions given for adenovirus serotype 5. Accordingly, the instant invention encompasses all adenoviral vectors partially deleted in E1 comprising basepairs corresponding to 1-450 (particularly, 342-450) and, preferably, 3511-3523 of a wild-type adenovirus serotype 5 (Ad5) nucleic acid sequence. Particularly preferred embodiments of the instant invention are those derived from adenoviruses like Ad5 which are classified in subgroup C (e.g., Ad2).

Vectors in accordance with the instant invention are at least partially deleted in E1. Preferably the E1 region is completely deleted or inactivated. Most preferably, the region deleted of E1 is within basepairs 451-3510. It is to be noted that the extended 5' and 3' regions of the disclosed vectors are believed to effectively reduce the size of the E1 deletion of previous constructs without overlapping any part of the E1A/E1B gene present in the cell line used, i.e., the PER.C6® cell line transefected with base pairs 459-3510. Overlap of adenoviral sequences is avoided because of the possibility of recombination. One of ordinary skill in the art can certainly appreciate that the instant invention can, therefore, be modified if a different cell line transfected with a different segment of adenovirus DNA is utilized. For purposes of exemplification, a 5' region of base pairs 1 to up to 449 is more appropriate if a cell line is transfected with adenoviral sequence from base pairs 450-3510. This holds true as well in the consideration of segments 3' to the E1 deletion.

Preferred embodiments of the instant invention possess an intact E3 region (i.e., an E3 gene capable of encoding a functional E3). Alternate embodiments have a partially deleted E3, an inactivated E3 region, or a sequence completely deleted of E3. Applicants have found, in accordance with the instant invention, that virus comprising the E3 gene were able to amplify more rapidly compared with virus not comprising an E3 gene; see Figure 6 wherein a diagnostic CsCl band corresponding to the E3+ virus

tested (5,665 bp) was present in greater amount compared with the diagnostic band of 3,010 bp corresponding to the E3- virus. These results were obtained following a virus competition study involving mixing equal MOI ratio (1:1) of adenovectors both comprising the E3 gene and not comprising the E3 gene. This increased amplification capacity of the E3+ adenovectors was subsequently confirmed with growth studies; see Table 4A, wherein the E3+ virus exhibit amplification ratios of 470, 420 and 320 as compared with the 115 and 40-50 of the E3- constructs.

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As stated above, vectors in accordance with the instant invention can accommodate up to approximately 4,850 base pairs of exogenous genetic material for an E3+ vector and approximately 7,500 base pairs for an E3- vector. Preferably, the insert brings the adenoviral vector as close as possible to a wild-type genomic size (e.g., for Ad5, 35,935 basepairs). It is well known that adenovirus amplifies best when they are close to their wild-type genomic size.

The genetic material can be inserted in an E1-parallel or an E1 anti-parallel orientation, as such is illustrated in Figure 7A, 7B, 7C and Figure 8A. Particularly preferred embodiments of the instant invention, have the insert in an E1-parallel orientation. Applicants have found, via competition experiments with plasmids containing transgenes in differing orientation (Figure 8A), that vector constructs with the foreign DNA insert in an E1-parallel orientation amplify better and actually outcompete E1-antiparallel-oriented transgenes. Viral DNA analysis of the mixtures at passage 3 and certainly at passage 6, showed a greater ratio of the virus carrying the transgene in the E1 parallel orientation as compared with the E1 anti-parallel version. By passage 10, the only viral species observed was the adenovector with the transgene in the E1 parallel orientation for both transgenes tested.

Adenoviral vectors in accordance with the instant invention are particularly well suited to effectuate expression of desired proteins, one example of which is an HIV protein, particularly an HIV full length gag protein. Exogenous genetic material encoding a protein of interest can exist in the form of an expression cassette. A gene expression cassette preferably comprises (a) a nucleic acid encoding a protein of interest; (b) a heterologous promoter operatively linked to the nucleic acid encoding the protein; and (c) a transcription terminator.

The transcriptional promoter is preferably recognized by an eukaryotic RNA polymerase. In a preferred embodiment, the promoter is a "strong" or "efficient" promoter. An example of a strong promoter is the immediate early human cytomegalovirus promoter (Chapman et al, 1991 *Nucl. Acids Res*19:3979-3986, which is incorporated by reference), preferably without intronic sequences. Most preferred

for use within the instant adenoviral vector is a human CMV promoter without intronic sequences, like intron A. Applicants have found that intron A, a portion of the human cytomegalovirus promoter (hCMV), constitutes a region of instability for adenoviral vectors. CMV without intron A has been found to effectuate (Examples 1-3) comparable expression capabilities in vitro when driving HIV gag expression and, furthermore, behaved equivalently to intron A-containing constructs in Balb/c mice in vivo with respect to their antibody and T-cell responses at both dosages of plasmid DNA tested (20 µg and 200 µg). Those skilled in the art will appreciate that any of a number of other known promoters, such as the strong immunoglobulin, or other eukaryotic gene promoters may also be used, including the EF1 alpha promoter, the murine CMV promoter, Rous sarcoma virus (RSV) promoter, SV40 early/late promoters and the beta-actin promoter.

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In preferred embodiments, the promoter may also comprise a regulatable sequence such as the Tet operator sequence. This would be extremely useful, for example, in cases where the gene products are effecting a result other than that desired and repression is sought.

The combination of the CMV promoter (devoid of the intron A region) with the BGH terminator is particularly preferred although other promoter/terminator combinations in the context of FG adenovirus may also be used.

Other embodiments incorporate a leader or signal peptide into the transgene. A preferred leader is that from the tissue-specific plasminogen activator protein, tPA. Examples include but are not limited to the various tPA-gag, tPA-pol and tPA-nef adenovirus-based vaccines disclosed throughout this specification.

In view of the improved adenovirus vectors described herein, an essential portion of the present invention are adenoviral-based HTV vaccines comprising said adenovirus backbones which may be administered to a mammalian host, preferably a human host, in either a prophylactic or therapeutic setting. The HTV vaccines of the present invention, whether administered alone or in combination regimens with other viral- or non-viral-based DNA vaccines, should elicit potent and broad cellular immune responses against HTV that will either lessen the likelihood of persistent virus infection and/or lead to the establishment of a clinically significant lowered virus load

subject to HTV infection or in combination with HAART therapy, mitigate the effects of previously established HIV infection (antiviral immunotherapy(ARI)). While any HIV antigen (e.g., gag, pol, nef, gp160, gp41, gp120, tat, rev, etc.) may be utilized in the herein described recombinant adenoviral vectors, preferred embodiments include the codon optimized p55 gag antigen (herein exemplified as MRKAd5gag), pol and nef. Sequences based on different Clades of HIV-1 are suitable for use in the instant invention, most preferred of which are Clade B and Clade C. Particularly preferred embodiments are those sequences (especially, codon-optimized sequences) based on concensus Clade B sequences. Preferred versions of the MRKAd5pol and MRKAd5nef series of adenoviral vaccines will encode modified versions of pol or nef, as discussed herein. Preferred embodiments of the MRKAd5HIV-1 vectors carrying HIV envelope genes and modifications thereof comprise the HIV codon-optimized env sequences of PCT International Applications PCT/US97/02294 and PCT/US97/10517, published August 28, 1997 (WO 97/31115) and December 24, 1997, respectively; both documents of which are hereby incorporated by reference.

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A most preferred aspect of the instant invention is the disclosed use of the adenoviral vector described above to effectuate expression of HIV gag. Sequences for many genes of many HIV strains are publicly available in GENBANK and primary, field isolates of HIV are available from the National Institute of Allergy and 20 Infectious Diseases (NIAID) which has contracted with Quality Biological (Gaithersburg, MD) to make these strains available. Strains are also available from the World Health Organization (WHO), Geneva Switzerland. It is preferred that the gag gene be from an HIV-1 strain (CAM-1; Myers et al, eds. "Human Retroviruses and AIDS: 1995, IIA3-IIA19, which is hereby incorporated by reference). This gene 25 closely resembles the consensus amino acid sequence for the clade B (North American/European) sequence. Therefore, it is within the purview of the skilled artisan to choose an appropriate nucleotide sequence which encodes a specific HIV gag antigen, or immunologically relevant portion thereof. As shown in Example 25, a clade B or clade C based p55 gag antigen will potentially be useful on a global scale. 30 As noted herein, the transgene of choice for insertion in to a DNA or MRKAd-based adenoviral vector of the present invention is a codon optimized version of p55 gag. Such a MRKAd5gag adenoviral vector is documented in Example 11 and is at least referred to herein as MRKAd5HIV-1gag. Of course, additional versions are contemplated, including but not limited to modifications such as promoter (e.g., mCMV for hCMV) and/or pA-terminations signal (SPA for bGH) switching, as well 35 as generating MRK Ad5 backbones with or without deletion of the Ad5 E3 gene.

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The present invention also relates a series of MRKAd5pol-based adenoviral vaccines which are shown herein to generate cellular immune responses subsequent to administration in mice and non-human primate studies. Several of the MRKAd5pol series are exemplified herein. One such adenoviral vector is referred to as MRKAd5hCMV-inact opt pol(E3+), which comprises the MRKAd5 backbone, the hCMV promoter (no intron A), an inactivated pol transgene, and contains the Ad5 E3 gene in the adenoviral backbone. A second exemplified pre-adenovirus plasmid and concomitant virus is referred to as MRKAd5hCMV-inact opt pol(E3-), which is identical to the former adenoviral vector except that the E3 is deleted. Both constructions contain a codon optimized, inactivated version of HIV-1 Pol, wherein at least the entire coding region is disclosed herein as SEQ ID NO:3 and the expressed protein is shown as SEQ ID NO:4 (see also Figure 17A-C and Table 1, which show targeted deletion for inactivated pol. This and other preferred codon optimized versions of HIV Pol as disclosed herein are essentially as described in U.S. Application Serial No. 09/745,221, filed December 21, 2000 and PCT International Application PCT/US00/34724, also filed December 21, 2000, both documents which are hereby incorporated by reference. As disclosed in the above-mentioned documents, the open reading frame for these codon-optimized HIV-1 Pol-based DNA vaccines are represented by codon optimized DNA molecules encoding codon optimized HIV-1 Pol (e.g. SEQ ID NO:2), codon optimized HIV-1 Pol fused to an amino terminal localized leader sequence (e.g. SEQ ID NO:6), and especially preferable, and exemplified by the MRKAd5-Pol construct in e.g., Example 19, biologically inactivated pol ("inact opt Pol"; e.g., SEQ ID NO:4) which is devoid of significant PR, RT, RNase or IN activity associated with wild type Pol. In addition, a construct related to SEQ ID NO:4 is contemplated which contains a leader peptide at the amino terminal region of the IA Pol protein. A specific construct is ligated within an appropriate DNA plasmid vector containing regulatory regions operatively linked to the respective HIV-1 Pol coding region, with or without a nucleotide sequence encoding a functional leader peptide. To this end, various HIV-1 Pol constructs disclosed herein relate to open reading frames for cloning to the enhanced first generation Ad vectors of the present invention (such a series of MRKAd5pol adenoviral vaccine vectors), including but not limited to wild type Pol (comprising the DNA molecule encoding WT opt Pol, as set forth in SEQ ID NO:2), tPA-opt WTPol, (comprising the DNA molecule encoding tPA Pol, as set forth in SEQ ID NO:6), inact 35 opt Pol (comprising the DNA molecule encoding IA Pol, as set forth in SEQ ID NO:4), and tPA-inact opt Pol, (comprising the DNA molecule encoding tPA-inact opt

Pol, as set forth in SEQ ID NO:8). The pol-based versions of enhanced first generation adenovirus vaccines elicit CTL and Th cellular immune responses upon administration to the host, including primates and especially humans. As noted in the above, an effect of the cellular immune-directed vaccines of the present invention should be a lower transmission rate to previously uninfected individuals and/or reduction in the levels of the viral loads within an infected individual, so as to prolong the asymptomatic phase of HIV-1 infection.

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The present invention further relates to a series of MRKAd5nef-based adenoviral vaccines which, similar to HIV gag and pol antigens, generate cellular immune responses subsequent to administration in mice and non-human primate 10 studies. The MRKAd5nef series are exemplified herein by utilizing the improved MRK adenoviral backbone in combination with modified versions of HIV nef. These exemplified MRKAd5nef vectors are as follows: (1) MRKAd5hCMVnef(G2A,LLAA) (E3+), which comprises the improved MRKAd5 backbone, a human CMV promoter an intact Ad5 E3 gene and a modified nef gene: (2) MRKAd5mCMV-15 nef(G2A,LLAA) (E3+), which is the same as (1) above but substituting a murine CMV promoter for a human CMV promoter; and (3) MRKAd5mCMV-tpanef(LLAA) (E3+), which is the same as (2) except that the nef transgene is tpanef(LLAA). Codon optimized versions of HIV-1 Nef and HIV-1 Nef modifications are essentially as described in U.S. Application Serial No. 09/738,782, filed December 15, 2000 and 20 PCT International Application PCT/US00/34162, also filed December 15, 2000, both documents which are hereby incorporated by reference. Particular embodiments of codon optimized Nef and Nef modifications relate to a DNA molecule encoding HIV-1 Nef from the HIV-1 jfrl isolate wherein the codons are optimized for expression in a mammalian system such as a human. The DNA molecule which encodes this protein 25 is disclosed herein as SEQ ID NO:9, while the expressed open reading frame is disclosed herein as SEQ ID NO:10. Another embodiment of Nef-based coding regions for use in the adenoviral vectors of the present invention comprise a codon optimized DNA molecule encoding a protein containing the human plasminogen activator (tpa) leader peptide fused with the NH₂-terminus of the HIV-1 Nef 30 polypeptide. The DNA molecule which encodes this protein is disclosed herein as SEQ ID NO:11, while the expressed open reading frame is disclosed herein as SEQ ID NO:12. Another modified Nef optimized coding region relates to a DNA molecule encoding optimized HIV-1 Nef wherein the open reading frame codes for modifications at the amino terminal myristylation site (Gly-2 to Ala-2) and 35 substitution of the Leu-174-Leu-175 dileucine motif to Ala-174-Ala-175, herein

described as opt nef (G2A, LLAA). The DNA molecule which encodes this protein is disclosed herein as SEQ ID NO:13, while the expressed open reading frame is disclosed herein as SEQ ID NO:14. MRKAd5nef vectors (1) MRKAd5hCMV-nef(G2A,LLAA) (E3+) and (2) MRKAd5mCMV-nef(G2A,LLAA) (E3+) contain this transgene. An additional embodiment relates to a DNA molecule encoding optimized HIV-1 Nef wherein the amino terminal myristylation site and dileucine motif have been deleted, as well as comprising a tPA leader peptide. This DNA molecule, opt tpanef (LLAA), comprises an open reading frame which encodes a Nef protein containing a tPA leader sequence fused to amino acid residue 6-216 of HIV-1 Nef (jfrl), wherein Leu-174 and Leu-175 are substituted with Ala-174 and Ala-175, herein referred to as opt tpanef (LLAA) is disclosed herein as SEQ ID NO:15, while the expressed open reading frame is disclosed herein as SEQ ID NO:16. The MRKAd5nef vector "MRKAd5mCMV-tpanef(LLAA) (E3+)" contains this transgene.

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Along with the improved MRKAd5gag adenovirus vaccine vector described herein, generation of a MRKAd5pol and MRKAd5nef adenovirus vector provide for enhanced HIV vaccine capabilities. Namely, the generation of this trio of adenoviral vaccine vectors, all shown to generate effective cellular immune responses subsequent to host administration, provide for the ability to administer these vaccine candidates not only alone, but preferably as part of a divalent (i.e., gag and nef, gag and pol, or pol and nef components) or a trivalent vaccine (i.e., gag, pol and nef components). Therefore, a preferred aspect of the present invention are vaccine formulations and associated methods of administration and concomitant generation of host cellular immune responses associated with formulating three separate series of MRKAd5based adenoviral vector vaccines. Of course, this MRKAd5 vaccine series based on distinct HIV antigens promotes expanded opportunities for formulation of a divalent or trivalent vaccine, or possibly administration of separate formulations of one or more monovalent or divalent formulations within a reasonable window of time. It is also within the scope of the present invention to embark on combined modality regimes which include multiple but distinct components from a specific antigen. An example, but certainly not a limitation, would be separate MRKAd5pol vectors, with one vaccine vector expressing wild type Pol (SEQ ID NO:2) and another MRKAd5pol vector expressing inactivated Pol (SEQ ID NO:6). Another example might be separate MRKAd5nef vectors, with one vaccine vector expressing the tPA/LLAA version of Nef (SEQ ID NO:16) and another MRKAd5nef vector expressing the G2A,LLAA modified version of Nef (SEQ ID NO:14). Therefore, the MRKAd5 adenoviral vectors of the present invention may be used in combination

with multiple, distinct HIV antigen classes. Each HIV antigen class is subject to sequence manipulation, thus providing for a multitude of potential vaccine combinations; and such combinations are within the scope of the present invention. The utilization of such combined modalities vaccine formulation and administration increase the probability of eliciting an even more potent cellular immune response when compared to inoculation with a single modality regimen.

The present invention also relates to application of a mono-, dual-, or trimodality administration regime of the MRKAd5gag, pol and nef adenoviral vaccine series in a prime/boost vaccination schedule. This prime/boost schedule may include any reasonable combination of the MRKAd5gag, pol and nef adenoviral vaccine series disclosed herein. In addition, a prime/boost regime may also involve other viral and/or non-viral DNA vaccines. A preferable addition to an adenoviral vaccine vector regime includes but is not limited to plasmid DNA vaccines, especially DNA plasmid vaccines that contain at least one of the codon optimized gag, pol and nef constructions, as disclosed herein.

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Therefore, one aspect of this invention is the administration of the adenoviral vector containing the optimized gag gene in a prime/boost regiment in conjunction with a plasmid DNA encoding gag. To distinguish this plasmid from the adenoviralcontaining shuttle plasmids used in the construction of an adenovirus vector, this plasmid will be referred to as a "vaccine plasmid" or "DNA plasmid vaccine". 20 Preferred vaccine plasmids for use in this administration protocol are disclosed in pending U.S. patent application 09/017,981, filed February 3, 1998 and WO98/34640, published August 13, 1998, both of which are hereby incorporated by reference. Briefly, the preferred vaccine plasmid is designated V1Jns-FLgag, which expresses the same codon-optimized gag gene as the adenoviral vectors of this invention (see 25 Figure 2 for the nucleotide sequence of the exemplified optimized codon version of full length p55 gag). The vaccine plasmid backbone, designated V1Ins contains the CMV immediate-early (IE) promoter and intron A, a bovine growth hormone-derived polyadenylation and transcription termination sequence as the gene expression regulatory elements, and a minimal pUC backbone; see Montgomery et al., 1993, 30 DNA Cell Biol. 12:777-783. The pUC sequence permits high levels of plasmid production in E. coli and has a neomycin resistance gene in place of an ampicillin resistance gene to provide selected growth in the presence of kanamycin. Alternatively, a vaccine plasmid which has the CMV promoter deleted of intron A can be used. Those of skill in the art will recognize that alternative vaccine plasmid 35

vectors may be easily substituted for these specific constructs, and this invention specifically envisions use of such alternative plasmid DNA vaccine vectors.

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Another aspect of the present invention is a prime/boost regimen which includes a vaccine plasmid which encodes an HIV pol antigen, preferably a codon optimized form of pol and also preferably a vaccine plasmid which comprises a nucleotide sequence which encodes a Pol antigen selected from the group of Pol antigens as shown in SEQ ID NOs: 2, 4, 6 and 8. The variety of potential DNA plasmid vaccines which encode various biologically active forms of HIV-1 Pol, wherein administration, intracellular delivery and expression of the HIV-1 Pol gene of interest elicits a host CTL and Th response. The preferred synthetic DNA molecules of the present invention encode codon optimized wild type Pol (without Pro activity) and various codon optimized inactivated HIV-1 Pol proteins. The HIV-1 pol open reading disclosed herein are especially preferred for pharmaceutical uses, especially for human administration as delivered via a recombinant adenoviral vaccine, especially an enhanced first generation recombinant adenoviral vaccine as described herein. Several embodiments of this portion of the invention are provided in detail below, namely DNA molecules which comprise a HIV-1 pol open reading frame, whether encoding full length pol or a modification or fusion as described herein, wherein the codon usage has been optimized for expression in a mammal, especially a human. Again, these DNA sequences are positioned appropriately within a recombinant adenoviral vector, such as the exemplified recombinant adenoviral vector described herein, so as to promote expression of the respective HIV-1 Pol gene of interest, and subsequent to administration, elicit a host CTL and Th response. Again, these preferred, but in no way limiting, pol genes are as disclosed herein and essentially as described in U.S. Application Serial No. 09/745,221, filed December 21, 2000 and PCT International Application PCT/US00/34724, also filed December 21, 2000, both documents which are hereby incorporated by reference.

A third series of vaccine plasmids which are useful in a combined modality and/or prime/boost regimen are vaccine plasmids which encode an HIV nef antigen or biologically and/or immunologically relevant modification thereof. As noted elsewhere, preferred vaccine plasmids contain a codon optimized form of nef and also preferably comprise a nucleotide sequence which encodes a Nef antigen selected from the group of Nef antigens as shown in SEQ ID NOs: 10, 12, 14 and 16. These preferred nef coding regions are disclosed herein, as well as being described in U.S. Application Serial No. 09/738,782, filed December 15, 2000 and PCT International

Application PCT/US00/34162, also filed December 15, 2000, both documents which are hereby incorporated by reference.

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Therefore, the adenoviral vaccines and plasmid DNA vaccines of this invention may be administered alone, or may be part of a prime and boost administration regimen. A mixed modality priming and booster inoculation scheme will result in an enhanced immune response, particularly is pre-existing anti-vector immune responses are present. This one aspect of this invention is a method of priming a subject with the plasmid vaccine by administering the plasmid vaccine at least one time, allowing a predetermined length of time to pass, and then boosting by administering the adenoviral vaccine. Multiple primings typically, 1-4, are usually employed, although more may be used. The length of time between priming and boost may typically vary from about four months to a year, but other time frames may be used. In experiments with rhesus monkeys, the animals were primed four times with plasmid vaccines, then were boosted 4 months later with the adenoviral vaccine. Their cellular immune response was notably higher than that of animals which had only received adenoviral vaccine. The use of a priming regimen may be particularly preferred in situations where a person has a pre-existing anti-adenovirus immune response.

Furthermore and in the alternative, multiple HIV-1 viral antigens, such as the MRKAd5 adenoviral vaccines disclosed herein, may be ligated into a proper shuttle 20 plasmid for generation of a pre-adenoviral plasmid comprising multiple open reading frames. For example a trivalent vector may comprise a gag-pol-nef fusion, in either a E3(-) or E3(+) background, preferably a E3 deleted backbone, or possible a "2+1" divalent vaccine, such as a gag-pol fusion (i.e., codon optimized p55 gag and inactivated optimized pol; Example 29 and Table 25) within the same MRKAd5 25 backbone, with each open reading frame being operatively linked to a distinct promoter and transcription termination sequence. Alternatively, the two open reading frames may be operatively linked to a single promoter, with the open reading frames operatively linked by an internal ribosome entry sequence (IRES), as disclosed in International Publication No. WO 95/24485, which is hereby incorporated by 30 reference. Figure 9 shows that the use of multiple promoters and termination sequences provide for similar growth properties, while Figure 28 shows that these MRKAd5gag-based vectors are also stable at least through passage 21. In the absence of the use of IRES-based technology, it is preferred that a distinct promoter be used to support each respective open reading frame, so as to best preserve vector stability. As 35 examples, and certainly not as limitations, potential multiple transgene vaccines may

include a three transgene vector such as hCMV-gagpol-bGHpA + mCMV-nef-SPA in an E3 deleted backbone or hCMV-gagpol-bGHpA + mCMV-nef-SPA(E3+). Potential "2+1" divalent vaccines of the present invention might be a hCMV-gagbGHpA + mCMV-nef-SPA in an E3+ backbone (vector #1) in combination with hCMV-pol-bGHpA in an E3+ backbone (vector #2), with all transgenes in the E1 parallel orientation. Fusion constructs other than the gag-pol fusion described above are also suitable for use in various divalent vaccine strategies and can be composed of any two HIV antigens fused to one another (e.g.,, nef-pol and gag-nef). These adenoviral compositions are, as above, preferably delivered along with an adenoviral composition comprising an additional HIV antigen in order to diversify the immune response generated upon administration. Therefore, a multivalent vaccine delivered in a single, or possible second, adenoviral vector is certainly contemplated as part of the present invention. Again, this mode of administration is another example of whereby an efficaceous adenovirus-based HIV-1 vaccine may be administered via a combined modality regime. It is important to note, however, that in terms of deciding on an insert for the disclosed adenoviral vectors, due consideration must be dedicated to the effective packaging limitations of the adenovirus vehicle. Adenovirus has been shown to exhibit an upper cloning capacity limit of approximately 105% of the wildtype Ad5 sequence.

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Regardless of the gene chosen for expression, it is preferred that the sequence be "optimized" for expression in a human cellular environment. A "triplet" codon of four possible nucleotide bases can exist in 64 variant forms. That these forms provide the message for only 20 different amino acids (as well as transcription initiation and termination) means that some amino acids can be coded for by more than one codon. Indeed, some amino acids have as many as six "redundant", alternative codons while some others have a single, required codon. For reasons not completely understood, alternative codons are not at all uniformly present in the endogenous DNA of differing types of cells and there appears to exist variable natural hierarchy or "preference" for certain codons in certain types of cells. As one example, the amino acid leucine is specified by any of six DNA codons including CTA, CTC, CTG, CTT, TTA, and TTG (which correspond, respectively, to the mRNA codons, CUA, CUC, CUG, CUU, UUA and UUG). Exhaustive analysis of genome codon frequencies for microorganisms has revealed endogenous DNA of E. coli most commonly contains the CTG leucine-specifying codon, while the DNA of yeasts and slime molds most commonly includes a TTA leucine-specifying codon. In view of this hierarchy, it is generally held that the likelihood of obtaining high levels of expression of a leucine-

rich polypeptide by an *E. coli* host will depend to some extent on the frequency of codon use. For example, a gene rich in TTA codons will in all probability be poorly expressed in *E. coli*, whereas a CTG rich gene will probably highly express the polypeptide. Similarly, when yeast cells are the projected transformation host cells for expression of a leucine-rich polypeptide, a preferred codon for use in an inserted DNA would be TTA.

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The implications of codon preference phenomena on recombinant DNA techniques are manifest, and the phenomenon may serve to explain many prior failures to achieve high expression levels of exogenous genes in successfully transformed host organisms—a less "preferred" codon may be repeatedly present in the inserted gene and the host cell machinery for expression may not operate as efficiently. This phenomenon suggests that synthetic genes which have been designed to include a projected host cell's preferred codons provide a preferred form of foreign genetic material for practice of recombinant DNA techniques. Thus, one aspect of this invention is an adenovirus vector or adenovirus vector in some combination with a vaccine plasmid where both specifically include a gene which is codon optimized for expression in a human cellular environment. As noted herein, a preferred gene for use in the instant invention is a codon-optimized HIV gene and, particularly, HIV gag, pol or nef.

Adenoviral vectors in accordance with the instant invention can be constructed using known techniques, such as those reviewed in Hitt et al, 1997 "Human Adenovirus Vectors for Gene Transfer into Mammalian Cells" Advances in Pharmacology 40:137-206, which is hereby incorporated by reference.

In constructing the adenoviral vectors of this invention, it is often convenient to insert them into a plasmid or shuttle vector. These techniques are known and described in Hitt et al., *supra*. This invention specifically includes both the adenovirus and the adenovirus when inserted into a shuttle plasmid.

Preferred shuttle vectors contain an adenoviral portion and a plasmid portion. The adenoviral portion is essentially the same as the adenovirus vector discussed supra, containing adenoviral sequences (with non-functional or deleted E1 and E3 regions) and the gene expression cassette, flanked by convenient restriction sites. The plasmid portion of the shuttle vector often contains an antibiotic resistance marker under transcriptional control of a prokaryotic promoter so that expression of the antibiotic does not occur in eukaryotic cells. Ampicillin resistance genes, neomycin resistance genes and other pharmaceutically acceptable antibiotic resistance markers may be used. To aid in the high level production of the polynucleotide by

fermentation in prokaryotic organisms, it is advantageous for the shuttle vector to contain a prokaryotic origin of replication and be of high copy number. A number of commercially available prokaryotic cloning vectors provide these benefits. It is desirable to remove non-essential DNA sequences. It is also desirable that the vectors not be able to replicate in eukaryotic cells. This minimizes the risk of integration of polynucleotide vaccine sequences into the recipients' genome. Tissue-specific promoters or enhancers may be used whenever it is desirable to limit expression of the polynucleotide to a particular tissue type.

In one embodiment of this invention, the pre-plasmids (e.g., pMRKAd5pol, pMRKAd5nef and pMRKAd5gag were generated by homologous recombination using the MRKHVE3 (and MRKHVO for the E3- version) backbones and the appropriate shuttle vector, as shown for pMRKAd5pol in Figure 22 and for pMRKAd5nef in Figure 23. The plasmid in linear form is capable of replication after entering the PER.C6[®] cells and virus is produced. The infected cells and media were harvested after viral replication was complete.

Viral vectors can be propagated in various E1 complementing cell lines, including the known cell lines 293 and PER.C6[®]. Both these cell lines express the adenoviral E1 gene product. PER.C6[®] is described in WO 97/00326 (published January 3, 1997) and issued U.S. Patent No. 6,033,908, both of which are hereby incorporated by reference. It is a primary human retinoblast cell line transduced with an E1 gene segment that complements the production of replication deficient (FG) adenovirus, but is designed to prevent generation of replication competent adenovirus by homologous recombination. Cells of particular interest have been stably transformed with a transgene that encodes the AD5E1A and E1B gene, like PER.C6[®], from 459 bp to 3510 bp inclusive. 293 cells are described in Graham et al., 1977 J. Gen. Virol 36:59-72, which is hereby incorporated by reference. As stated above, consideration must be given to the adenoviral sequences present in the complementing cell line used. It is important that the sequences not overlap with that present in the vector if the possibility of recombination is to be minimized.

It has been found that vectors generated in accordance with the above description are more effective in inducing an immune response and, thus, constitute very promising vaccine candidates. More particularly, it has been found that first generation adenoviral vectors in accordance with the above description carrying a codon-optimized HIV gag gene, regulated with a strong heterologous promoter can be used as human anti-HIV vaccines, and are capable of inducing immune responses.

Standard techniques of molecular biology for preparing and purifying DNA constructs enable the preparation of the DNA immunogens of this invention.

A vaccine composition comprising an adenoviral vector in accordance with the instant invention may contain physiologically acceptable components, such as buffer, normal saline or phosphate buffered saline, sucrose, other salts and polysorbate. One preferred formulation has: 2.5-10 mM TRIS buffer, preferably about 5 mM TRIS buffer; 25-100 mM NaCl, preferably about 75 mM NaCl; 2.5-10% sucrose, preferably about 5% sucrose; 0.01 -2 mM MgCl₂; and 0.001%-0.01% polysorbate 80 (plant derived). The pH should range from about 7.0-9.0, preferably about 8.0. One skilled in the art will appreciate that other conventional vaccine excipients may also be used it make the formulation. The preferred formulation contains 5mM TRIS, 75 mM NaCl, 5% sucrose, 1mM MgCl₂, 0.005% polysorbate 80 at pH 8.0 This has a pH and divalent cation composition which is near the optimum for Ad5 stability and minimizes the potential for adsorption of virus to a glass surface. It does not cause tissue irritation upon intramuscular injection. It is preferably frozen until use.

The amount of adenoviral particles in the vaccine composition to be introduced into a vaccine recipient will depend on the strength of the transcriptional and translational promoters used and on the immunogenicity of the expressed gene product. In general, an immunologically or prophylactically effective dose of 1×10^7 to 1×10^{12} particles and preferably about 1×10^{10} to 1×10^{11} particles is administered directly into muscle tissue. Subcutaneous injection, intradermal introduction, impression through the skin, and other modes of administration such as intraperitoneal, intravenous, or inhalation delivery are also contemplated. It is also contemplated that booster vaccinations are to be provided. Following vaccination with HIV adenoviral vector, boosting with a subsequent HIV adenoviral vector and/or plasmid may be desirable. Parenteral administration, such as intravenous, intramuscular, subcutaneous or other means of administration of interleukin-12 protein, concurrently with or subsequent to parenteral introduction of the vaccine compositions of this invention is also advantageous.

The adenoviral vector and/or vaccine plasmids of this invention polynucleotide may be unassociated with any proteins, adjuvants or other agents which impact on the recipients' immune system. In this case, it is desirable for the vector to be in a physiologically acceptable solution, such as, but not limited to, sterile saline or sterile buffered saline. Alternatively, the vector may be associated with an adjuvant known in the art to boost immune responses (i.e., a "biologically effective"

adjuvant), such as a protein or other carrier. Vaccine plasmids of this invention may, for instance, be delivered in saline (e.g., PBS) with or without an adjuvant. Preferred adjuvants are Alum or CRL1005 Block Copolymer. Agents which assist in the cellular uptake of DNA, such as, but not limited to, calcium ions, may also be used to advantage. These agents are generally referred to herein as transfection facilitating reagents and pharmaceutically acceptable carriers. Techniques for coating microprojectiles coated with polynucleotide are known in the art and are also useful in connection with this invention.

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This invention also includes a prime and boost regimen wherein a first adenoviral vector is administered, then a booster dose is given. The booster dose may be repeated at selected time intervals. Alternatively, a preferred inoculation scheme comprises priming with a first adenovirus serotype and then boosting with a second adenovirus serotype. More preferably, the inoculation scheme comprises priming with a first adenovirus serotype and then boosting with a second adenovirus serotype, wherein the first and second adenovirus serotypes are classified within separate subgroups of adenoviruses. The above prime/boost schemes are particularly preferred in those situations where a preexisting immunity is identified to the adenoviral vector of choice. In this type of scheme, the individual or population of individuals is primed with an adenovirus of a serotype other than that to which the preexisting immunity is identified. This enables the first adenovirus to effectuate sufficient expression of the transgene while evading existing immunity to the second adenovirus (the boosting adenovirus) and, further, allows for the subsequent delivery of the transgene via the boosting adenovirus to be more effective. Adenovirus serotype 5 is one example of a virus to which such a scheme might be desirable. In accordance with this invention, therefore, one might decide to prime with a non-group C adenovirus (e.g., Ad12, a group A adenovirus, Ad24, a group D adenovirus, or Ad35, a group B adenovirus) to evade anti-Ad5 immunity and then boost with Ad5, a group C adenovirus. Another preferred embodiment involves administration of a different adenovirus (including non-human adenovirus) vaccine followed by administration of the adenoviral vaccines disclosed. In the alternative, a viral antigen of interest can be first delivered via a viral vaccine other than an adenovirus-based vaccine, and then followed with the adenoviral vaccine disclosed. Alternative viral vaccines include but are not limited to pox virus and venezuelan equine encephilitis virus.

A large body of human and animal data supports the importance of cellular immune responses, especially CTL in controlling (or eliminating) HIV infection. In humans, very high levels of CTL develop following primary infection and correlate

with the control of viremia. Several small groups of individuals have been described who are repeatedly exposed to HIV by remain uninfected; CTL has been noted in several of these cohorts. In the SIV model of HIV infection, CTL similarly develops following primary infection, and it has been demonstrated that addition of anti-CD8 monoclonal antibody abrogated this control of infection and leads to disease progression. This invention uses adenoviral vaccines alone or in combination with plasmid vaccines to induce CTL.

The following non-limiting Examples are presented to better illustrate the invention.

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EXAMPLE 1

Removal of the Intron A Portion of the hCMV Promoter

GMP grade pVIInsHIVgag was used as the starting material to amplify the hCMV promoter. PVIInsHIVgag is a plasmid comprising the CMV immediate-early (IE) promoter and intron A, a full-length codon-optimized HIV gag gene, a bovine growth hormone-derived polyadenylation and transcriptional termination sequence, and a minimal pUC backbone; see Montgomery et al., supra for a description of the plasmid backbone. The amplification was performed with primers suitably positioned to flank the hCMV promoter. A 5' primer was placed upstream of the Msc1 site of the hCMV promoter and a 3' primer (designed to contain the BgIII recognition sequence) was placed 3' of the hCMV promoter. The resulting PCR product (using high fidelity Taq polymerase) which encompassed the entire hCMV promoter (minus intron A) was cloned into TOPO PCR blunt vector and then removed by double digestion with Msc1 and BgIII. This fragment was then cloned back into the original GMP grade pV1InsHIVgag plasmid from which the original promoter, intron A, and the gag gene were removed following Msc1 and BgIII digestion. This ligation reaction resulted in the construction of a hCMV promoter (minus intron A) + bGHpA

The FLgag gene was excised from pV1JnsHIVgag using BgIII digestion and the 1,526 bp gene was gel purified and cloned into pV1JnsCMV(no intron) at the BgIII site. Colonies were screened using Sma1 restriction enzymes to identify clones that carried the Flgag gene in the correct orientation. This plasmid, designated pV1JnsCMV(no intron)-FLgag-bGHpA, was fully sequenced to confirm sequence integrity.

expression cassette within the original pV1JnsHIVgag vector backbone. This vector

is designated pVIInsCMV(no intron).

Two additional transgenes were also constructed. The plasmid, pV1JnsCMV(no intron)-FLgag-SPA, is identical to pV1JnsCMV(no intron)-FLgag-bGHpA except that the bovine growth hormone polyadenylation signal has been replaced with a short synthetic polyA signal (SPA) of 50 nucleotides in length. The sequence of the SPA is as shown, with the essential components (poly(A) site, (GT)_n, and (T)_n; respectively) underlined:

<u>AATAAA</u>AGATCTTTATTTTCATTAGATCT<u>GTGTG TTGGTTTTTTGTGTG</u> (SEQ ID NO:18).

The plasmid, pV1Jns-mCMV-FLgag-bGHpA, is identical to the pV1JnsCMV(no intron)-FLgag-bGHpA except that the hCMV promoter has been removed and replaced with the murine CMV (mCMV) promoter.

Figure 3 diagrammatically shows the new transgene constructs in comparison with the original transgene.

15 EXAMPLE 2

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Gag Expression Assay for Modified Gag Transgenes

Gag Elisa was performed on culture supernatants obtained from transient tissue culture transfection experiments in which the two new hCMV-containing plasmid constructs, pV1JnsCMV(no intron)-FLgag-bGHpA and pV1JnsCMV(no intron)-FLgag-SPA, both devoid of intron A, were compared to pV1JnsHIVgag which, as noted above possesses the intron A as part of the hCMV promoter. Table 2 below shows the *in vitro* gag expression data of the new gag plasmids compared with the GMP grade original plasmid. The results displayed in Table 2 show that both of the new hCMV gag plasmid constructs have expression capacities comparable to the original plasmid construct which contains the intron A portion of the hCMV promoter.

Table 2: In vitro DNA transfection of original and new plasmid HIV-1 gag constructs.

Plasmid	μg gag/10e6 COS cells/5μg DNA/48 hr
HIVFL-gagPR9901ª	10.8
PVIIns-hCMV-FLgag-bGHpAb	16.6
pV1Jns-hCMV-FLgag-SPA ^{b,c}	12.0

^a GMP grade pV1 Jns-hCMV intronA-FL gag-bGHpA.

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EXAMPLE 3

Rodent (Balb/c) Study for Modified gag Transgenes
A rodent study was performed on the two new plasmid constructs
described above – pV1JnsCMV(no intron)-FLgag-bGHpA and pV1JnsCMV(no
intron)-FLgag-SPA - in order to compare them with the construct described above
possessing the intron A portion of the CMV promoter, pV1JnsHIVgag. Gag antibody
and Elispot responses (described in PCT International Application No.
PCT/US00/18332 (WO 01/02607) filed July 3, 2000, claiming priority to U.S.
Provisional Application Serial No. 60/142,631, filed July 6, 1999 and U.S.
Application Serial No. 60/148,981, filed August 13, 1999, all three applications which
are hereby incorporated by reference) were measured. The results displayed in Table
3 below, show that the new plasmid constructs behaved equivalently to the original
construct in Balb/c mice with respect to their antibody and T-cell responses at both
dosages of plasmid DNA tested, 20 μg and 200 μg.

⁵ b New plasmid constructions that have the intron A portion removed from the hCMV promoter.

^c In this construct the bGH terminator has been replaced with the short synthetic polyadenylation signal (SPA)

EXAMPLE 4

Table 3: HIV191: Immunogenicity of V1Jns-gag under different promoter and termination control elements.

DNA ^a	Dose,	Anti-p24 Titers			SFC/10^6 Cells			
	ug ^b	(3 Wk PD1) ^c			(4 Wk PD1) ^d			
Promoter/terminator		GMT	+SE	-SE	Media	gag197-205	p24	
HIVFL-gagPR9901	200	12800	4652	3412	2(2)	129(19)	30(11)	
(GMP grade)	20	5572	1574	1227	0	56(9)	25(6)	
pV1Jns-hCMV-	200	11143	2831	2257	0	98(5)	12(6)	
FL-gag-bGHpA	20	7352	2808	2032		73(9)	11(6)	
pV1Jns-hCMV-	200	16890	5815	4326	1(1)	94(4)	26(7)	
FL-gag-SPA	20	5971	5361	2825	0	85(17)	38(10)	
Naīve	o	123	50	36	0	0	0	

^ein PBS

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Construction of the Modified Shuttle Vector - "MRKpdelE1 Shuttle"

The modifications to the original Ad5 shuttle vector (pdelE1sp1A; a vector comprising Ad5 sequences from basepairs 1-341 and 3524-5798, with a multiple cloning region between nucleotides 341 and 3524 of Ad5, included the following three manipulations carried out in sequential cloning steps as follows:

- (1) The left ITR region was extended to include the *Pac1* site at the junction between the vector backbone and the adenovirus left ITR sequences. This allow for easier manipulations using the bacterial homologous recombination system.
- 10 (2) The packaging region was extended to include sequences of the wild-type (WT) adenovirus from 342 bp to 450 bp inclusive.
 - (3) The area downstream of pIX was extended 13 nucleotides (i.e., nucleotides 3511-3523 inclusive).
- These modifications (Figure 4) effectively reduced the size of the E1 deletion without overlapping with any part of the E1A/E1B gene present in the transformed PER.C6[®] cell line. All manipulations were performed by modifying the Ad shuttle vector pdelE1sp1A.

Once the modifications were made to the shuttle vector, the changes were incorporated into the original Ad5 adenovector backbones (pAdHVO and pAdHVE3) by bacterial homologous recombination using *E. coli* BJ5183 chemically competent cells.

bi.m. Injections into both quads, 50 µL per quad

cn=10:GMT, geometric mean titer; SE, standard, error

dn=5, pooled spleens; mean of triplicate wells and standard, deviation, in parentheses;

EXAMPLE 5

Construction of Modified Adenovector Backbones (E3+ and E3-)

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The original adenovectors pAdHVO (comprising all Ad5 sequences except those nucleotides encompassing the E1 and E3 regions) and pADHVE3 (comprising all Ad5 sequences except those nucleotides encompassing the E1 region), were each reconstructed so that they contained the modifications to the E1 region. This was accomplished by digesting the newly modified shuttle vector (MRKpdelE1 shuttle) with Pac1 and BstZ1101 and isolating the 2,734 bp fragment which corresponds to the adenovirus sequence. This fragment was co-transformed with DNA from either Cla1 linearized pAdHVO (E3- adenovector) or Cla1 linearized pAdHVE3 (E3+adenovector) into E. coli BJ5183 competent cells. At least two colonies from each transformation were selected and grown in Terrific™ broth for 6-8 hours until turbidity was reached. DNA was extracted from each cell pellet and then transformed into E. coli XL1 competent cells. One colony from each transformation was selected and grown for plasmid DNA purification. The plasmid was analyzed by restriction digestions to identify correct clones. The modified adenovectors were designated MRKpAdHVO (E3- plasmid) and MRKpAdHVE3 (E3+ plasmid). Virus from these new adenovectors (MRKHVO and MRKHVE3, respectively) as well as the old version of the adenovectors were generated in the PER.C6® cell lines to accommodate the following series of viral competition experiments. In addition, the multiple cloning site of the original shuttle vector contained ClaI, BamHI, Xho I, EcoRV, HindIII, Sal I, and Bgl II sites. This MCS was replaced with a new MCS containing Not I, Cla I, EcoRV and Asc I sites. This new MCS has been transferred to the MRKpAdHVO and MRKpAdHVE3 pre-plasmids along with the modification made to the packaging region and pIX gene.

EXAMPLE 6

Analysis of the Effect of the Packaging Signal Extension

To study the effects of the modifications made to the E1 deletion region, the viruses obtained from the original backbone (pAdHVE3) and the new backbone (MRKpAdHVE3) were mixed together in equal MOI ratios (1:1 and 5:5) and passaged through several rounds; see Figure 5, Expt.#1. Both of the viruses in the experiment contained the E3 gene intact and did not contain a transgene. The only difference between the two viruses was within the region of the E1 deletion. Following the coinfection of the viruses at P1 (passage 1), the mixtures were propagated through an additional 4 passages at which time the cells were harvested

and the virus extracted and purified by CsCl banding. The viral DNA was extracted and digested with *Hind*III and the digestion products were then radioactively labeled. For the controls, the respective pre-plasmids (pAdHVE3 ("OLD E3+"); MRKpAdHVE3 ("NEW E3+")) were also digested with *Hind*III (and *Pac*1 to remove the vector backbone) and subsequently labeled with [³³P]dATP. The radioactively labeled digestion products were subjected to gel electrophoresis and the gel was dried down onto Whatman paper before being exposed to autoradiographic film. Figure 6 clearly shows that the new adenovirus which has the addition made to the packaging signal region has a growth advantage compared with the original adenovirus. In the experiments performed (at either ratio tested), only the digestion bands pertaining to the newly modified virus were present. The diagnostic band of size 3,206 (from the new virus) was clearly present. However, there was no evidence of the diagnostic band of size 2,737 bp expected from the original virus.

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EXAMPLE 7

Analysis of the Effect of the E3 Gene

The second set of the virus competition study involved mixing equal MOI ratio (1:1) of the newly modified viruses, that obtained from MRKpAdHVO and MRKpAdHVE3 (Figure 5, Expt. #2). In this set, both viruses had the new modifications made to the E1 deletion. The first virus (that from MRKpAdHVO) does not contain an E3 gene. The second virus (that from MRKpAdHVE3) does contain the E3 gene. Neither of the viruses contain a transgene. Following coinfection of the viruses, the mixtures were propagated through an additional 4 passages at which time the cells were harvested and the total virus extracted and purified by CsCl banding. The viral DNA was extracted and digested with HindIII and the digestion products were then radioactively labeled. For the controls, the respective pre-plasmids MRKpAdHVO ("NEW E3-"); MRKpAdHVE3 ("NEW E3+") were also digested with Hind (and Pac1 to remove the vector backbone) and then labeled with [33P]dATP. The radioactively labeled digestion products were subjected to gel electrophoresis and the gel was dried down onto Whatman paper before being exposed to autoradiographic film. Figure 6 shows the results of the viral DNA analysis of the E3+ virus and E3- virus mixing experiment. The diagnostic band corresponding to the E3+ virus (5,665 bp) was present in greater amount compared with the diagnostic band of 3,010 bp corresponding to the E3- virus. This indicates that the virus that contains the E3 gene is able to amplify more rapidly

compared with the virus that does not contain an E3 gene. This increased amplification capacity has been confirmed by growth studies; see Table 4 below.

EXAMPLE 8

Construction of the new shuttle vector containing modified gag transgene – "MRKpdelE1-CMV(no intron)-FLgag-bGHpA"

The modified plasmid pV1JnsCMV(no intron)-FLgag-bGHpA was digested with *Msc*1 overnight and then digested with *Sfi*1 for 2 hours at 50°C. The DNA was then treated with Mungbean nuclease for 30 mins at 30°C. The DNA mixture was desalted using the Qiaex II kit and then Klenow treated for 30 mins at 37°C to fully blunt the ends of the transgene fragment. The 2,559 bp transgene fragment was then gel purified. The modified shuttle vector (MRKpdelE1 shuttle) was linearized by digestion with EcoRV, treated with calf intestinal phosphatase and the resulting 6,479 bp fragment was then gel purified. The two purified fragments were then ligated together and several dozen clones were screened to check for insertion of the transgene within the shuttle vector. Diagnostic restriction digestion was performed to identify those clones carrying the transgene in the E1 parallel and E1 anti-parallel orientation. This strategy was followed to clone in the other gag transgenes in the MRKpdelE1 shuttle vector.

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EXAMPLE 9

Construction of the MRK FG Adenovectors

The shuttle vector containing the HIV-1 gag transgene in the E1 parallel orientation, MRKpdelE1-CMV(no intron)-FLgag-bGHpA, was digested with Pac1. The reaction mixture was digested with BsfZ171. The 5,291 bp fragment was purified by gel extraction. The MRKpAdHVE3 plasmid was digested with Cla1 overnight at 37°C and gel purified. About 100 ng of the 5,290 bp shuttle +transgene fragment and ~100 ng of linearized MRKpAdHVE3 DNA were co-transformed into E. coli BJ5183 chemically competent cells. Several clones were selected and grown in 2 ml Terrific™ broth for 6-8 hours, until turbidity was reached. The total DNA from the cell pellet was purified using Qiagen alkaline lysis and phenol chloroform method. The DNA was precipitated with isopropanol and resuspended in 20 μl dH₂0. A 2 μl aliquot of this DNA was transformed into E. coli XL-1 competent cells. A single colony from each separate transformation was selected and grown overnight in 3 ml LB +100 μg/ml ampicillin. The DNA was isolated using Qiagen columns. A positive clone was identified by digestion with the restriction enzyme BstEII which cleaves

within the gag gene as well as the plasmid backbone. The pre-plasmid clone is designated MRKpAdHVE3+CMV(no intron)-FLgag-bGHpA and is 37,498 bp in size. This strategy was followed to generate E3- and E3+ versions of each of the other gag transgene constructions in both E1 parallel and E1 anti-parallel versions. Figures 7A, 7B and 7C show the various combinations of adenovectors constructed.

EXAMPLE 10

Plasmid Competition Studies

A series of plasmid competition studies was carried out. Briefly, the screening of the various combinations of new constructs was performed by mixing equal amounts of each of two competing plasmids. In the experiment shown in Figure 8A, plasmids containing the same transgene but in different orientations were mixed together to create a "competition" between the two plasmids. The aim was to look at the effects of transgene orientation. In the experiment shown in Figure 8B, plasmids containing different polyadenylation signals (but in the same orientation) were mixed together in equal amounts. The aim was to assess effects of polyA signals. Following the initial transfection, the virus was passaged through ten rounds and the viral DNA analyzed by radioactive restriction analysis.

Analysis of the viral species from the plasmid mixing experiment (Figure 8A) showed that adenovectors which had the transgene inserted in the E1 parallel orientation amplified better and were able to out-compete the adenovirus which had the transgene inserted in the E1 anti-parallel orientation. Viral DNA analysis of the mixtures at passage 3 and certainly at passage 6, showed a greater ratio of the virus carrying the transgene in the E1 parallel orientation compared with the E1 antiparallel version. By passage 10, the only viral species observed was the adenovector with the transgene in the E1 parallel orientation for both transgenes tested (hCMV(no intron)-FLgag-bGHpA and hCMV(no intron)-FLgag-SPA).

Analysis of the viral species from the plasmid mixing experiment #2 (Figure 8B) at passages 3 and 6 showed that the polyadenylation signals tested (bGHpA and SPA) did not have an effect on the growth of the virus. Even at passage 10 the two viral species in the mixture were still present in equal amounts.

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EXAMPLE 11

Virus generation of an enhanced adenoviral construct - "MRK Ad5 HIV-1gag"

The results obtained from the competition study allowed us to make the following conclusions: (1) The packaging signal extension is beneficial; (2) Presence of E3 does enhance viral growth; (3) E1 parallel orientation is recommended; and (4) PolyA signals have no effect on the growth of the adenovirus.

MRK Ad5 HIV-1 gag exhibited the most desirable results. This construct contains the hCMV(no intron)-FLgag-bGHpA transgene inserted into the new E3+ adenovector backbone, MRKpAdHVE3, in the E1 parallel orientation. We have designated this adenovector MRK Ad5 HIV-1 gag. This construct was prepared as outlined below:

The pre-plasmid MRKpAdHVE3+CMV(no intron)-FLgag-bGHpA was digested was Pac1 to release the vector backbone and 3.3 µg was transfected by calcium phosphate method (Amersham Pharmacia Biotech.) in a 6 cm dish containing PER.C6® cells at ~60% confluence. Once CPE was reached (7-10 days), the culture was freeze/thawed three times and the cell debris pelleted. 1 ml of this cell lysate was used to infect into a 6 cm dish containing PER.C6® cells at 80-90% confluence. Once CPE was reached, the culture was freeze/thawed three times and the cell debris pelleted. The cell lysate was then used to infect a 15 cm dish containing PER.C6[®] cells at 80-90% confluence. This infection procedure was continued and expanded at passage 6. The virus was then extracted from the cell pellet by CsCl method. Two bandings were performed (3-gradient CsCl followed by a continuous CsCl gradient). Following the second banding, the virus was dialyzed in A105 buffer. Viral DNA was extracted using pronase treatment followed by phenol chloroform. The viral DNA was then digested with *Hind*III and radioactively labeled with [33P]dATP. Following gel electrophoresis to separate the digestion products the gel was dried down on Whatman paper and then subjected to autoradiography. The digestion products were compared with the digestion products from the pre-plasmid (that had been digested with Pac1/HindIII prior to labeling). The expected sizes were observed, indicating that the virus had been successfully rescued. This strategy was used to rescue virus from each of the various adenovector plasmid constructs prepared.

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EXAMPLE 12

Stability Analyses

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To determine whether the various adenovector constructs (e.g., MRK Ad5 HIV-1 gag) show genetic stability, the viruses were each passaged continually. The viral DNA was analyzed at passages 3, 6 and 10. Each virus maintained its correct genetic structure. In addition, the stability of the MRK Ad5 HIV-1 gag was analyzed under propagation conditions similar to that performed in large scale production. For this analysis, the transfections of MRK Ad5 HIV-1 gag as well as three other adenoviral vectors were repeated and the virus was purified at P3. The three other adenovectors were as follows: (1) that comprising hCMV(no intron)-Flgag with a bGHpA terminator in an E3- adenovector backbone; (2) that comprising hCMV(no intron)-Flgag with a SPA termination signal in an E3+ adenovector backbone, and that comprising a mCMV-Flgag with a bGHpA terminator in an E3+ adenovector backbone. All of the vectors have the transgene inserted in the E1 parallel orientation. Viral DNA was analyzed by radioactive restriction analysis to confirm that it was correct before being delivered to fermentation cell culture for continued passaging in serum-free media. At P5 each of the four viruses were purified and the viral DNA extracted for analysis by the restriction digestion and radiolabeling procedure. This virus has subsequently been used in a series of studies (in vitro gag expression in COS cells, rodent study and rhesus monkey study) as will be described below. The viruses from P5 are shown in Figure 9.

The passaging under serum-free conditions was continued for the MRKHVE3 (transgene-less, obtained from MRKpAdHVE3 pre-plasmid) and the MRKAd5HIV-1gag (obtained from MRKpAdHVE3+CMV(no intron)-FLgag-bGHpA pre-plasmid) viruses. Figure 10 shows viral DNA analysis by radioactive restriction digestion at passage 11 for MRKHVE3, MRKAd5HIV-1gagE3-, and passage 11 and 12 for MRKAd5HIV-1gag. Aside from the first lane which is the DNA marker lane, the next three lanes are virus from the pre-plasmid controls (controls based on the original virus) - MRKpAdHVE3 (also referred to as "pMRKHVE3"), MRKpAdHVE3+CMV(no intron)-FLgag-bGHpA, and pMRKAd5gag(E3-), respectively. As seen in Figure 10, each of the viral DNA samples show the expected bands with no extraneous bands showing. This signifies that there are no major variant adenovirus species present that can be detected by autoradiography.

Figure 11 shows the results of viral competition study between MRKHVE3 and MRKAd5HIV-1gag. These viruses were mixed together at equal MOI (140 viral

particles each; 280 vp total) at passage 6 and continued to be passaged until P11. Aside from the first lane which is the DNA marker lane, the next two lanes are the pre-plasmid controls obtained from MRKpAdHVE3 and MRKpAdHVE3+CMV(no intron)-FLgag-bGHpA. The next two lanes are the viral DNA from the starting viral material at passage six. The last two lanes are the competition studies performed in duplicate. The data in Figure 11 shows the effect the gag transgene in culture. Growth of a MRKAd5gag virus was compared with growth of a "transgene-less" MRKHVE3. These two viruses were infected at the same MOI (i.e. 140 vp each) at passage 6 and then passaged through to passage 11 and the viral pool was analyzed by radioactive restriction analysis. The data shows that one virus did not out compete the other. Therefore, the gag transgene did not show obvious signs of toxicity to the adenovirus.

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Analysis by *Hind*III digestion shows that each virus specie is present in approximately equal amounts. As above, there does not appear to be signs of any extraneous bands. Figure 12 shows higher passage numbers for MRKAd5HIV-1gag grown under serum-containing conditions. The genome integrity again has been maintained and there is no evidence of rearrangements, even at the highest passage level (P21).

Each of the four vectors shown in Figure 9 were analyzed for amplification capacity. Table 4 below shows the QPA analysis used in the estimation of viral amplification ratios at P4. The determination of the amplification ratio for the original HIV-1 gag construct is based on the clinical lot at P12. It has been shown that amplification rates increases with higher passage number for the original virus. The reason for this observation is due to the emergence of variants which exhibit increased growth rates compared to the intact adenovector. With continued passaging of the original Ad gag vector, the level of variants increases and hence amplification rates increase also.

The MRK Ad5 HIV-1 gag virus has also been continually passaged under process conditions (i.e., serum-free media). Viral DNA extracted from passages 11 and 12 show no evidence of rearrangement.

Table 4:
Amplification Ratios Based on AEX and QPA Analysis of Virus Amplification from Passage 3 to Passage 4.

Ad gag construct	Amplification Ratio
MRKAd5gag	470
HCMV-Flgag-bGHpA [E3-]	115
HCMV-Flgag-SPA [E3+]	320
mCMV-FLgag-bGHpA [E3+]	420
Original construct *	40 - 50

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EXAMPLE 13 Analytical Evaluation of the enhanced Ad5 Constructs

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To study the effects of the transgene and the E3 gene on virus amplification, the enhanced adenoviral vector, MRK Ad5 HIV-1 gag, along with its transgene-less version (MRKpAdHVE3) and its E3- version (MRK Ad5 HIV-1 gag E3-), was studied for several passages under serum-free conditions. Table 5A shows the amplification ratios determined for passages P3 to P8 for MRK Ad5 HIV-1 gag. Within a certain MOI range, it has been determined that the virus output is directly proportional to the virus input. Therefore, the greater the number of virus particles per cell at infection, the greater the virus amount produced. Viral amplification ratios,

on the other hand, are inversely proportional to the virus input. The lower the virus

input, the greater the amplification ratio.

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Table 5B shows the amplification rates of the new E3+ vector backbone MRKpAdHVE3. It has a significantly lower rate of amplification compared with the gag transgene containing version. This may be contributed to the larger size MRK Ad5 HIV-1 gag since it contains the transgene. This inclusion of the transgene brings the size of the adenovirus closer to the size of a wild type Ad5 virus. It is well known that adenoviruses amplify best when they are at close to their wild type genomic size.

^{*} This estimation is based on the clinical lot growth characteristics at Passage 12.

Wild type Ad5 is 35,935 bp. The MRKpAdHVE3 is 32, 905 bp in length. The enhanced adenovector MRK Ad5 HIV-1 gag is 35,453bp (See Figure 14 for vector map; see also Figure 15A-X show the complete pre-adenoviral vector sequence, which includes an additional 2,021 bp of the vector backbone).

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Table 5C shows the amplification rates of the new E3- gag containing virus MRK Ad5 HIV-1 gag E3-. Once again, this virus shows lower growth rate than the enhanced adenoviral vector. This may be attributed to the decreased sized of this virus (due to the E3 gene deletion) compared with wild type Ad5. The MRK Ad5 HIV-1 gag E3- virus is 32,810 bp in length. This can be compared with the wild type Ad5 which is 35,935 bp and MRK Ad5 HIV-1 gag which is 35,453 bp in length.

Table 5A: Amplification ratios determined by AEX and QPA for MRKAd5gag over several continuous passaging in serum free media. Following P5, two replicate samples were taken (rep-1 and rep-2) and analyzed.

MRKAd5gag rep1

	Xv (10° cells/n	il), Viability (%)	Harvest Time	Cell Passage	Titer	Ther	OPA	Ratio	Amplification	AEX
	Infection	Harvest	hal	Number	10" vp/ml culture	10° vp/cell	10° TCiD _{er} /mi	AEX:QPA	Ratto	Internal Control
P4	1,49, 81%	0.58, 50%	44	46	8.7	5.9	1.72	50	470 (MOI = 125)	
P5	1.38, 93%	0.66, 47%	48	49	6.7	4.9	1.38	49	170	
P8	1.04, 94%	0.68, 77%	47	48	5.8	5.6	1.42	41	200	
P7	1.50, 84%	0.96, 61%	49.5	50	3.9	1.4	Q.97	40	50	
P7	1.09, 97%	0.78, 59%	50	52	5.2	4.7	1.70	81	170	
P8	1.03, 94%	0.86, 64%	47.5	54	9.0	8.7	1.10	82	310	
P9	0,89,95%	0.99, 73%	47.5	56	4,4	4.9	1.03	43	175	3.12 2.84
P10	1.09, 91%	1.08, 66%	47.5	68	8.0	2.8	1.16	26	100	2.70 2.60
P11	1.19, 88%	0.88, 65%	47	60	3.6	3.0	1.15	31	110	2.70 2.70
P12	0.98, 91%	0.85, 63%	47.5	47	5.4	5.5	1.20	45	200	2.86 2.60
P13	1.00, 88%	0.70, 67%	49	49	5.6	5.8	1.11	52	210	3.18 3.18
P14	1,94, 92%	0.88, 67%	46	53	8.6	4.4			160	3.28 3.27
P15	0.97, 96%	0.64, 56%	47	47	6.9	7.1			250	3.12 2.91

Table 5B: Amplification ratios determined by AEX and QPA for MRKHVE3 over several continuous passaging in serum free media. MRKHVE3 is the new vector backbone which does NOT carry a transgene.

MRKHVE3

	Xv (10° cells/n	ni), Viability (%)	Harvest Time	Cell Passage	Titer	Titer	QPA	Ratio	Amplification	AEX
	Infection	Harvest	h,p.l.	Number	10 ^{to} vp/ml cutture	10° vp/ceti	10° TCID _{so} /ml	AEX:QPA	Ratio	Internal Control
P4	1.10, 97%	1.28, 79%	49	54	4.1	3.8	1.70	25	300 (MOI = 125)	-
P5	0.82, 89%	1.18, 77%	47	. 48	4.3	4.7	1.24	35	170	
P6	1,55, 88%	1,26, 76%	49.5	50	1.2	0.8	0.56	21	30	
P6	1.09, 97%	1.11, 81%	49	52	4.0	3.6	1.16	34	130	
P7	1.17, 91%	1,22, 91%	47.5	54	3.7	3.2	0.50	74	110	
P8	0.98, 88%	1,41, 83%	48	56	21	2.1	0.47	45	75	3.12 2.84
P9	1.20, 89%	1.26, 81%	47.5	58	8.0	0.7	0.29	28	25	2.70 2.60
P10	0.99, 82%	1.55, 85%	47	60	2.3	2.3	0.43	53	80	270 270
P11	1.07, 96%	1.25, 83%	48	47	2.7	2.5	0.41	66	90	2.66 2.60
P12	0.80, 91%	1.14, 60%	49,5	49	5.9	7.4	0.48	123	260	3.18 3.18
P13	1.96, 95%	1.14, 85%	45.6	53	5.8	3.0			110	3.28 3.27
P14	0.97, 96%	1.03, 98%	48.5	47	9.4	9.7			350	3.12 2.91
P15	0.87, 99%	0.97, 59%	49.5	49	5.3	6.1			218	2.78 2.52

Table 5C. Amplification ratios determined by AEX and QPA for MRKAd5gag(E3-) over several continuous passaging in serum free media. This construct is identical to the MRKAd5gag construct except that this version is DELETED of the E3 gene.

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MRKAd5qaq(E3-)

	Xv (10° cells/n	ni), Viability (%) Harvest	Harvest Time	Cell Passage Number	Titer 10 ¹⁰ vp/ml culture	Titer 10° vp/celi	QPA 10° TCID ₆₀ /ml	Ratio	Amplification	AEX
			h.p.l.					AEX:QPA	Ratio	Internal Contro
P4	1.62, 77%	1.12, 62%	47.5	46	2.0	1.2	0.92	20	100 (MOI=125)	
P5	1.16, 92%	0.62, 43%	49	49	3.3	2.9	0.99	34	100	
P6	1.71, 86%	0.20, 10%	49	50	4.7	2.7	1.70	28	100	
P6	1.09, 97%	0.63, 54%	49.5	52	5.4	5.0	1.76	31	180	
P7	1.17, 91%	0.98, 72%	47.50	54	7.1	6.1	0.67	106	220	
P8	0.98, 88%	0.77, 48%	48	56	3.1	3.2	0.66	47	115	3.12 2.84
P9	1.20, 89%	1.03, 72%	48	58	1.8	1.5	0.57	32	55	2.70 2.60
P10	0.99, 82%	0,80, 62%	46.5	60	3.2	3.2	0.68	47	115	2.70 2.70
P11	1.07, 96%	0.98, 70%	48.5	47	5.9	5.5	0.68	87	200	2.86 2.60
P12	0.80, 91%	0.67, 59%	50	49	5.1	6.4	0.72	71	230	3.18 3.18
P13	1.96, 95%	0.91, 59%	45.5	53	7.4	3.8			135	3.28 3.27
P14	0.97, 96%	0.81, 74%	48	47	6.8	7.0		-	250	3.12 2.91
P15	0.87, 99%	0.84, 56%	49	49	4.8	5.5			196	2.78 2.52

EXAMPLE 14

Gag Expression Analysis of the Novel Constructs

In vitro gag analysis of the MRK Ad5 HIV-1 gag and the original HIV-gag vectors (research and clinical lot) show comparable gag expression. The clinical lot shows only a slightly reduced gag expression level. The most noticeable difference is with the mCMV vector. This vector shows roughly 3 fold lower expression levels compared with the other vectors tested (which all contain hCMV promoters). The mCMV-FLgag with bGHpA assay was performed three times using different propagation and purification lots and it consistently exhibited weaker gag expression.

EXAMPLE 15

Evaluation of MRK Ad5 HIV-1 gag and Other gag-Containing Adenovectors in Balb/c Mice

Cohorts of 10 balb/c mice were vaccinated intramuscularly with escalating doses of MRK Ad5 HIV-1 gag, and the research and clinical lots of original Ad5HIV-1gag. Serum samples were collected 3 weeks post dose 1 and analyzed by anti-p24 sandwich ELISA.

Anti-p24 titers in mice that received MRK Ad5 HIV-1 gag (107 and 109 vp(viral particle) doses) were comparable (Figure 13) to those of the research lot of Ad5HIV-1 gag, for which much of the early rhesus data were generated on. These titers were also comparable when E3 is deleted (MRKAd5hCMVgagbGHpA(E3-)) or SPA is substituted for bGHpA terminator (MRKAd5 hCMV-gag-SPA (E3+)) or murine CMV promoter is used in place of hCMV (MRKAd5 mCMV-gag-bGHpA (E3+)) in the MRKAd5 backbone.

The results shown in Table 7 indicate that the three other vectors (in addition to the preferred vector, MRK Ad5 HIV-1 gag, are also capable of inducing strong anti-gag antibody responses in mice. Interestingly enough, while the mCMV-FLgag construct containing bGHpA and E3+ in an E1 parallel orientation showed lowest gag expression in the COS cell *in vitro* infection (Table 6) in comparison with the other vectors tested, it generated the greatest anti-gag antibody response this *in vivo* Balb/c study. Table 7 also shows a dose response in anti-gag antibody production in both the research and the clinical lot. As expected, the clinical lot shows reduced anti-gag antibody induction at each dosage level compared to the same dosage used for the research lot.

Table 6: In vitro analysis for gag expression in COS cells by Elisa assay.

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Viral Vectors ^a	μg gag/4.8x10e5 COS/10e8 parts/48hr
MRKAd5gag ^b	1.40
Clinical lot Ad5gag ^c	1.28
Research lot Ad5gag ^d	1.32
MCMVFL-gagbGHpA ^e	0.42

^a A_{260nm} absorbance readings taken for viral particle determinations.

^b MRKAd5gag was produced in serum free conditions and purified at P5.

^c Clinical lot# Ad5gagFN0001

d Research Ad5FLgag lot# 6399

^emCMVFL-gagbGHpA was produced in serum free conditions and purified at P5.

Table 7: mHIV020 Anti-p24 Ab Titers in Balb/c mice (n=10) vaccinated with various Adgag constructs and lots (3 week post dose1).

Group ID	Vaccine	Dose (vp)	GMT	SE upper	SE lower
4	^a MRKAd5gag	10^7	25600	5877	4780
2	mnkAdagag n	10^9	409600	94028	76473
3	hCMV FL-gag bGHpA [E3-] →	10^7	7352	2077	1620
4	в	10^9	235253	59767	47659
5	hCMV FL-gag SPA [E3+] →	10^7	12800	9905	236
6	•	10^9	310419	99181	75165
7	^b mCMV FL-gag bGHpA [E3+] →	10^7	44572	23504	15389
8	•	10^9	941014	239068	190636
9	^c hCMV FL-gag bGHpA [E3-] ←	10^7	3676	934	745
10	•	10^9	117627	17491	15227
11	research lot hCMV intronA FL-gag bGHpA [E3-] <-		528	262	175
12	n	10^7 10^8	14703 58813	5274 14942	3882 11915
14	"	10^9	204800	53232	42250
15	clinical lot hCMVintronA FL-gag bGHpA [E3-] <-	10^6	230	82	61
16		10^7	4222	3405	1138
17 18	"	10^8 10^9	19401 89144	3939 25187	3274 19639
19	Naĭve	none	93	7	6

*2x50 µL i.m. (quad) injections/animal

P.I.s: Youil, Chen, Casimiro Vaccination: T. Toner, Q. Su

Assay: M. Chen

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^aThe structure of MRKAd5gag is: hCMVFL-gagbGHpA [E3+] → The <u>same lot</u> of MRKAd5gag used in this rodent study was used in the Rhesus monkey study (Tables 7 and 8).

^bThe same lot of mCMVFL-gagbGHpA[E3+] used in the *in vitro* study (Table 6) ws used here.

EXAMPLE 16

Comparison of Humoral and Cellular Responses Towards the Original Ad-gag Construct with the New MRK Ad5 HIV-1 gag in Rhesus Monkeys

Cohorts of 3 rhesus monkeys were vaccinated intramuscularly with MRK Ad5 HIV-1 gag or the clinical Ad5gag bulk at two doses, 10^{11} vp and 10^9 vp.

Immunizations were conducted at week 0, 4, and 25. Serum and PBMC samples were collected at selected time points. The serum sample were assayed for anti-p24 Ab titers (using competitive based assay) and the PBMCs for antigen-specific IFN-gamma secretion following overnight stimulation with gag 20-mer peptide pool (via ELISpot assay).

The results shown in Table 8 indicate comparable responses with respect to the generation of anti-gag antibodies. The frequencies of gag-specific T cells in

^cThis construct was designed by Volker Sandig. It contains a shorter version of the hCMV promoter than that used in the MRK constructs. The adenovector backbone is identical to the original backbone used in the original Adgag vector. Expression at 10e7 dose from this vector is 7 fold lower then the same dose of the MRKAd5gag and 4 fold lower than the research lot.

peripheral blood assummarized in Table 9 demonstrate a strong cellular immune response generated after a single dose with the new construct MRK Ad5 HIV-1 gag. The responses are also boostable with second dose of the same vector. The vector is also able to induce CD8+ T cell responses (as evident by remaining spot counts after CD4+ depletion of PBMCs) which are responsible for cytotoxic activity.

Table 8 Anti-p24 antibody titers (in mMU/mL) in rhesus macaques immunized with

gag-expressing adenovectors (Protocol HIV203).

gag-expressing adenovector				1 394. 30	2010. 2.4	1141-00	1111.00	144-00
Vaccine	Pre_	Wk4	Wk8	Wk 12	Wk 16_	Wk 20	Wk 25	Wk 28
MRKAd5gag ^a , 10^11 vp								
97N010	<10	118	5528_	11523	7062	21997	ND	51593
97N116	<10	62	772_	1447	1562	2174	ND	20029
98X007	<10	66.	3353	6156	6845	3719	ND_	24031
MRKAd5gog, 10^9 vp								
97N120	<10	51	204	318	366	482	ND	6550
97N144	<10	18	118	274	706	888	ND	7136
98X008	<10	15	444	386	996	1072	ND	12851
Ad5gag ^b , Clinical Lot, 10^11 vp								
97X001	<10	87	2579_	4718	7174	7250	ND	69226
97N146	<10	72	3604_	7380	7526	18906	ND	60283
98X009	<10	78	4183	3946	3124	6956	ND_	26226
Ad5gag, Clinical Lot, 10^9 vp								
97N020	<10	<10	143	371	390	1821	ND	17177
97X003	<10	<10	39	93	156	596_	ND	2053
98X012	<10	81	342	717	956	1558	ND	11861
MRKAd5gag (hCMV, bGHpA, E3+)		l	1	ļ		<u> </u>		<u> </u>
^b original Adagag vector (hCMV/Intro	n A bGHp	A, E3-), lot	#FN0001_		<u> </u>	ļ		ļ
ND, not determined		<u> </u>		<u> </u>	<u> </u>	<u> </u>		1

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Table 9. Number of gag-specific T cells per million peripheral blood mononuclear cells (PBMCs) in rhesus monkeys immunized with gag-expressing adenovectors. Also included are those frequencies in PBMCs depleted of CD4⁺ T cells.

Vaccination	Monkey ID				Wk	1=1	Wk	I=1	5 Wk				Wk
T=0,4,25 wks		Media ^a	Gog H ^b	Media	Gog H	Media	Gog H	Media	Gog H	Media	Gog H	Media	Goog H
MRKActions	97N010	6	89	0	395	0	1058	0	1174	3	775 74	4	1074 594
1041 VP				1 1	609			4	305	ĭ			408
		1	676	1		ŏ	593		0.0	o	184	Ŏ	666
	98X007	10	579	0	1304	3	2193	1	2118	3	1588	0	2113
	98X007(CD4-)	20	965			0	2675			0	1656	٥	1278
MRKAc5gcg	97N120	5	275	1	249	4	141	4	119	9	206	4	219
10/9 vp				١.	400			١ ,	۸.,			!	219 373
				l B	438			3	250				625
				Ι.	1000			١,	673				735
	98X008(CD4-)	14	696	\	1030	ŏ	1175	"	1 5/3	ŏ	391	4	848
AdScoop dinical lat	97X001	0	261	1	485	0	817	0	1220b	1	894	0	1858
10/11 vp	97X001(CD4-)	10	283		1								1123
	97N146	3		1	465			1	1272				1785
		٥		١.				١ _	١				971
		-		3	339			0	876				1748
	98X009(CD4-)	0	/3			<u> </u>	333				225	L	044
Adagag diniad lat	97N020	3	30	1	101	0	66	0	36	0	26	00	41 16
IU-9 VP				, s	194			١,	38		38		81
				ľ			1	'	1		4	١ŏ	19
			95	3	54	Ιĭ	34	0	18	Ĭŏ	20	i	121
	98X012(CD4-)	ii	70			Ó	11			0	В	0	41
Naive	96R041	6	8	1	1	0	0	0	0	0	0	1 24	0
	T=0,4,25 wks MRKAd5gag 10/41 vp MRKAd5gag 10/9 vp Ad5gag dinical lat 10/41 vp	T=0,4,25 wks MRKAd5gcg 97N010 10/41 vp 97N010(CD4-) 97N116 97N116 97N116 97N116 97N116 97N120 10/9 vp 97N120 97N120 97N120 97N120 97N120(CD4-) 98X003 98X003(CD4-) Ad5gcg dinical lot 97X001 10/41 vp 97X001(CD4-) 98X009 98X009(CD4-) Ad5gcg dinical lot 97X001 10/41 vp 97X001(CD4-) 98X009 98X009(CD4-) Ad5gcg dinical lot 97X001 10/9 vp 97X003 97N020(CD4-) 98X012 98X012(CD4-) 98X012 98X012(CD4-)	T=0,4,25 w/s Media* MRKAc5gcg 97ND10 6 10^11 vp 97N010(CD4-) 4 97N116 1 97N116(CD4-) 11 98X007 10 98X007(CD4-) 20 MRKAc5gcg 97N120 5 10^9 vp 97N120(CD4-) 11 97N144 3 97N144(CD4-) 6 98X008 4 98X008(CD4-) 14 Ac5gcg clinical lot 97X001 0 10^11 vp 97X001(CD4-) 10 97X014(CD4-) 6 98X009 0 98X009 0 98X009(CD4-) 0 Ac5gcg clinical lot 97N020 3 97N120(CD4-) 0 Ac5gcg clinical lot 97N020 3 97N020(CD4-) 10 97X003 4 97X003 5 97X003 5 97X003 5 97X003 5 97X003 5 97X003 10 97X003 10 97X003 10 97X003 10 97X003 10 97X003 10 97X004 10 97X004 10 97X004 10 97X005 11	T=0,4,25 wks	T=0,4,25 wks	T=0.4.25 w/s Media¹ Gog H³ Media¹ Gog H³ MRKAc5gcq 10^41 vp 97ND10 97N010(CD4-) 98X007 6 89 0 395 97N116 97N116(CD4-) 10 98X007 (CD4-) 10 98X007 (CD4-) 98X007 (CD4-) 10 97N120(CD4-) 10 97N144 (CD4-) 97N144 (CD4-) 98X008 4 368 1 1090 1 249 Ac5gcq clinical lot 10^11 vp 97X001 97X014(CD4-) 97X001 0 283 97N146 3 150 1 465 97N146 (CD4-) 97X001 0 93 3 339 1 485 Ac5gcq clinical lot 10^12 vp 97X001 0 281 1 485 97N146 (CD4-) 97X003 4 68 5 134 Ac5gcq clinical lot 10^9 vp 97X002 3 3 30 1 101 Ac6gcq clinical lot 10^9 vp 97X003 3 30 1 101 Ac6gcq clinical lot 10^9 vp 97X002 3 3 30 1 101 97X003 4 68 5 134 97X003 4 68 5 134 97X003 4 68 5 134 97X001 5 98X012 5 95 3 54 98X012 5 95 3 54 98X012 5 95 3 54	T=0.4.25 w/s	T=0,4,25 w/s	T=0,4,25 wks	T=0.4.25 w/s MRKAc5gcg	T=0,4,25 wks	T=0,4,25 wks	T=0,4,25 wkg Media Gog H Media Gog H

Based on either 4x10/5 or 2x10/5 cells per well (depending on spot density)

ND, not determined

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Prock or no peptide control

The adenovectors described herein and, particularly, MRK Ad5 HIV-1 gag, represent very promising HIV-gag adenovectors with respect to their enhanced growth characteristics in both serum and, more importantly, in serum-free media conditions. In comparison with the current HIV-1 gag adenovector construct, MRK Ad5 HIV-1 gag shows a 5-10 fold increased amplification rate. We have shown that it is genetically stable at passage 21. This construct is able to generate significant cellular immune responses *in vivo* even at a relatively low dose of 10^9 vp. The potency of the MRKAd5gag construct is comparable to, if not better than the original HIV-1gag vector as shown in this rhesus monkey study.

EXAMPLE 17 CODON OPTIMIZED HIV-1 POL AND CODON OPTIMIZED HIV-1 POL MODIFICATIONS

The open reading frames for the various synthetic *pol* genes disclosed herein comprise coding sequences for the reverse transcriptase (or RT which consists of a polymerase and RNase H activity) and integrase (IN). The protein sequence is based

Pool of 20-capepitides overlapping by 10 caland encompassing the page sequence

on that of Hxb2r, a clonal isolate of IIIB; this sequence has been shown to be closest to the consensus clade B sequence with only 16 nonidentical residues out of 848 (Korber, et al., 1998, Human retroviruses and AIDS, Los Alamos National Laboratory, Los Alamos, New Mexico). The skilled artisan will understand after review of this specification that any available HTV-1 or HIV-2 strain provides a potential template for the generation of HIV pol DNA vaccine constructs disclosed herein. It is further noted that the protease gene is excluded from the DNA vaccine constructs of the present invention to insure safety from any residual protease activity in spite of mutational inactivation. The design of the gene sequences for both wildtype (wt-pol) and inactivated pol (IA-pol) incorporates the use of human preferred ("humanized") codons for each amino acid residue in the sequence in order to maximize in vivo mammalian expression (Lathe, 1985, J. Mol. Biol. 183:1-12). As can be discerned by inspecting the codon usage in SEQ ID NOs: 1, 3, 5 and 7, the following codon usage for mammalian optimization is preferred: Met (ATG), Gly (GGC), Lys (AAG), Trp (TGG), Ser (TCC), Arg (AGG), Val (GTG), Pro (CCC), Thr (ACC), Glu (GAG); Leu (CTG), His (CAC), Ile (ATC), Asn (AAC), Cys (TGC), Ala (GCC), Gln (CAG), Phe (TTC) and Tyr (TAC). For an additional discussion relating to mammalian (human) codon optimization, see WO 97/31115 (PCT/US97/02294), which, as noted elsewhere in this specification, is hereby incorporated by reference. It is intended that the skilled artisan may use alternative versions of codon optimization or may omit this step when generating HIV pol vaccine constructs within the scope of the present invention. Therefore, the present invention also relates to non-codon optimized versions of DNA molecules and associated recombinant adenoviral HIV vaccines which encode the various wild type and modified forms of the HIV Pol protein disclosed herein. However, codon optimization of these constructs is a preferred embodiment of this invention.

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A particular embodiment of this portion of the invention comprisies codon optimized nucleotide sequences which encode wt-pol DNA constructs (herein, "wt-pol" or "wt-pol (codon optimized))" wherein DNA sequences encoding the protease (PR) activity are deleted, leaving codon optimized "wild type" sequences which encode RT (reverse transcriptase and RNase H activity) and IN integrase activity. A DNA molecule which encodes this protein is disclosed herein as SEQ ID NO:1, the open reading frame being contained from an initiating Met residue at nucleotides 10-12 to a termination codon from nucleotides 2560-2562. SEQ ID NO:1 is as follows:

AGATCTACCA TGGCCCCCAT CTCCCCCATT GAGACTGTGC CTGTGAAGCT GAAGCCTGGC

ATGGATGGCC CCAAGGTGAA GCAGTGGCCC CTGACTGAGG AGAAGATCAA GGCCCTGGTG

	GAAATCTGCA	CTGAGATGGA	GAAGGAGGC	AAAATCTCCA	AGATTGGCCC	CGAGAACCCC
	TACAACACCC	CTGTGTTTGC	CATCAAGAAG	AAGGACTCCA	CCAAGTGGAG	GAAGCTGGTG
	GACTTCAGGG	AGCTGAACAA	GAGGACCCAG	GACTTCTGGG	AGGTGCAGCT	GGGCATCCCC
	CACCCCGCTG	GCCTGAAGAA	GAAGAAGTCT	GTGACTGTGC	TGGATGTGGG	GGATGCCTAC
5	TTCTCTGTGC	CCCTGGATGA	GGACTTCAGG	AAGTACACTG	CCTTCACCAT	CCCCTCCATC
	AACAATGAGA	CCCCTGGCAT	CAGGTACCAG	TACAATGTGC	TGCCCCAGGG	CTGGAAGGGC
	TCCCCTGCCA	TCTTCCAGTC	CTCCATGACC	AAGATCCTGG	AGCCCTTCAG	GAAGCAGAAC
	CCTGACATTG	TGATCTACCA	GTACATGGAT	GACCTGTATG	TGGGCTCTGA	CCTGGAGATT
	GGGCAGCACA	GGACCAAGAT	TGAGGAGCTG	AGGCAGCACC	TGCTGAGGTG	GGGCCTGACC
10	ACCCCTGACA	AGAAGCACCA	GAAGGAGCCC	CCCTTCCTGT	GGATGGGCTA	TGAGCTGCAC
	CCCGACAAGT	GGACTGTGCA	GCCCATTGTG	CTGCCTGAGA	AGGACTCCTG	GACTGTGAAT
	GACATCCAGA	AGCTGGTGGG	CAAGCTGAAC	TGGGCCTCCC	AAATCTACCC	TGGCATCAAG
	GTGAGGCAGC	TGTGCAAGCT	GCTGAGGGGC	ACCAAGGCCC	TGACTGAGGT	GATCCCCCTG
	ACTGAGGAGG	CTGAGCTGGA	GCTGGCTGAG	AACAGGGAGA	TCCTGAAGGA	GCCTGTGCAT
15	GGGGTGTACT	ATGACCCCTC	CAAGGACCTG	ATTGCTGAGA	TCCAGAAGCA	GGGCCAGGGC
	CAGTGGACCT	ACCAAATCTA	CCAGGAGCCC	TTCAAGAACC	TGAAGACTGG	CAAGTATGCC
	AGGATGAGGG	GGGCCCACAC	CAATGATGTG	AAGCAGCTGA	CTGAGGCTGT	GCAGAAGATC
	ACCACTGAGT	CCATTGTGAT	CTGGGGCAAG	ACCCCCAAGT	TCAAGCTGCC	CATCCAGAAG
	GAGACCTGGG	AGACCTGGTG	GACTGAGTAC	TGGCAGGCCA	CCTGGATCCC	TGAGTGGGAG
20	TTTGTGAACA	CCCCCCCT	GGTGAAGCTG	TGGTACCAGC	TGGAGAAGGA	GCCCATTGTG
	GGGGCTGAGA	CCTTCTATGT	GGATGGGGCT	GCCAACAGGG	AGACCAAGCT	GGGCAAGGCT
	GGCTATGTGA	CCAACAGGGG	CAGGCAGAAG	GTGGTGACCC	TGACTGACAC	CACCAACCAG
	AAGACTGAGC	TCCAGGCCAT	CTACCTGGCC	CTCCAGGACT	CTGGCCTGGA	GGTGAACATT
	GTGACTGACT	CCCAGTATGC	CCTGGGCATC	ATCCAGGCCC	AGCCTGATCA	GTCTGAGTCT
25	GAGCTGGTGA	ACCAGATCAT	TGAGCAGCTG	ATCAAGAAGG	AGAAGGTGTA	CCTGGCCTGG
	GTGCCTGCCC	ACAAGGGCAT	TGGGGGCAAT	GAGCAGGTGG	ACAAGCTGGT	GTCTGCTGGC
	ATCAGGAAGG	TGCTGTTCCT	GGATGGCATT	GACAAGGCCC	AGGATGAGCA	TGAGAAGTAC
	CACTCCAACT	GGAGGGCTAT	GGCCTCTGAC	TTCAACCTGC	CCCCTGTGGT	GGCTAAGGAG
	ATTGTGGCCT	CCTGTGACAA	GTGCCAGCTG	AAGGGGGAGG	CCATGCATGG	GCAGGTGGAC
30	TGCTCCCCTG	GCATCTGGCA	GCTGGACTGC	ACCCACCTGG	AGGGCAAGGT	GATCCTGGTG
	GCTGTGCATG	TGGCCTCCGG	CTACATTGAG	GCTGAGGTGA	TCCCTGCTGA	GACAGGCCAG
	GAGACTGCCT	ACTTCCTGCT	GAAGCTGGCT	GGCAGGTGGC	CTGTGAAGAC	CATCCACACT
	GACAATGGCT	CCAACTTCAC	TGGGGCCACA	GTGAGGGCTG	CCTGCTGGTG	GGCTGGCATC
	AAGCAGGAGT	TTGGCATCCC	CTACAACCCC	CAGTCCCAGG	GGGTGGTGGA	GTCCATGAAC
35	AAGGAGCTGA	AGAAGATCAT	TGGGCAGGTG	AGGGACCAGG	CTGAGCACCT	GAAGACAGCT
	GTGCAGATGG	CTGTGTTCAT	CCACAACTTC	AAGAGGAAGG	GGGGCATCGG	GGGCTACTCC

GCTGGGGAGA GGATTGTGGA CATCATTGCC ACAGACATCC AGACCAAGGA GCTCCAGAAG
CAGATCACCA AGATCCAGAA CTTCAGGGTG TACTACAGGG ACTCCAGGAA CCCCCTGTGG
AAGGGCCCTG CCAAGCTGCT GTGGAAGGGG GAGGGGGCTG TGGTGATCCA GGACAACTCT
GACATCAAGG TGGTGCCCAG GAGGAAGGCC AAGATCATCA GGGACTATGG CAAGCAGATG
GCTGGGGATG ACTGTGTGGC CTCCAGGCAG GATGAGGACT AAAGCCCGGG CAGATCT (SEQ
ID NO:1).

The open reading frame of the wild type pol construct disclosed as SEQ ID NO:1 contains 850 amino acids, disclosed herein as SEQ ID NO:2, as follows: Met Ala Pro Ile Ser Pro Ile Glu Thr Val Pro Val Lys Leu Lys Pro Gly Met Asp Gly Pro Lys Val Lys Gln Trp Pro Leu Thr Glu Glu Lys 10 Ile Lys Ala Leu Val Glu Ile Cys Thr Glu Met Glu Lys Glu Gly Lys Ile Ser Lys Ile Gly Pro Glu Asn Pro Tyr Asn Thr Pro Val Phe Ala Ile Lys Lys Lys Asp Ser Thr Lys Trp Arg Lys Leu Val Asp Phe Arg Glu Leu Asn Lys Arg Thr Gln Asp Phe Trp Glu Val Gln Leu Gly Ile Pro His Pro Ala Gly Leu Lys Lys Lys Ser Val Thr Val Leu Asp 15 Val Gly Asp Ala Tyr Phe Ser Val Pro Leu Asp Glu Asp Phe Arg Lys Tyr Thr Ala Phe Thr Ile Pro Ser Ile Asn Asn Glu Thr Pro Gly Ile Arg Tyr Gln Tyr Asn Val Leu Pro Gln Gly Trp Lys Gly Ser Pro Ala Ile Phe Gln Ser Ser Met Thr Lys Ile Leu Glu Pro Phe Arg Lys Gln 20 Asn Pro Asp Ile Val Ile Tyr Gln Tyr Met Asp Asp Leu Tyr Val Gly Ser Asp Leu Glu Ile Gly Gln His Arg Thr Lys Ile Glu Glu Leu Arg Gln His Leu Leu Arg Trp Gly Leu Thr Thr Pro Asp Lys Lys His Gln Lys Glu Pro Pro Phe Leu Trp Met Gly Tyr Glu Leu His Pro Asp Lys Trp Thr Val Gln Pro Ile Val Leu Pro Glu Lys Asp Ser Trp Thr Val Asn Asp Ile Gln Lys Leu Val Gly Lys Leu Asn Trp Ala Ser Gln Ile 25 Tyr Pro Gly Ile Lys Val Arg Gln Leu Cys Lys Leu Leu Arg Gly Thr Lys Ala Leu Thr Glu Val Ile Pro Leu Thr Glu Glu Ala Glu Leu Glu Leu Ala Glu Asn Arg Glu Ile Leu Lys Glu Pro Val His Gly Val Tyr Tyr Asp Pro Ser Lys Asp Leu Ile Ala Glu Ile Gln Lys Gln Gly Gln Gly Gln Trp Thr Tyr Gln Ile Tyr Gln Glu Pro Phe Lys Asn Leu Lys 30 Thr Gly Lys Tyr Ala Arg Met Arg Gly Ala His Thr Asn Asp Val Lys Gln Leu Thr Glu Ala Val Gln Lys Ile Thr Thr Glu Ser Ile Val Ile Trp Gly Lys Thr Pro Lys Phe Lys Leu Pro Ile Gln Lys Glu Thr Trp Glu Thr Trp Trp Thr Glu Tyr Trp Gln Ala Thr Trp Ile Pro Glu Trp 35 Glu Phe Val Asn Thr Pro Pro Leu Val Lys Leu Trp Tyr Gln Leu Glu Lys Glu Pro Ile Val Gly Ala Glu Thr Phe Tyr Val Asp Gly Ala Ala

Asn Arg Glu Thr Lys Leu Gly Lys Ala Gly Tyr Val Thr Asn Arg Gly Arg Gln Lys Val Val Thr Leu Thr Asp Thr Thr Asn Gln Lys Thr Glu Leu Gln Ala Ile Tyr Leu Ala Leu Gln Asp Ser Gly Leu Glu Val Asn Ile Val Thr Asp Ser Gln Tyr Ala Leu Gly Ile Ile Gln Ala Gln Pro Asp Gln Ser Glu Ser Glu Leu Val Asn Gln Ile Ile Glu Gln Leu Ile Lys Lys Glu Lys Val Tyr Leu Ala Trp Val Pro Ala His Lys Gly Ile Gly Gly Asn Glu Gln Val Asp Lys Leu Val Ser Ala Gly Ile Arg Lys Val Leu Phe Leu Asp Gly Ile Asp Lys Ala Gln Asp Glu His Glu Lys Tyr His Ser Asn Trp Arg Ala Met Ala Ser Asp Phe Asn Leu Pro Pro Val Val Ala Lys Glu Ile Val Ala Ser Cys Asp Lys Cys Gln Leu Lys 10 Gly Glu Ala Met His Gly Gln Val Asp Cys Ser Pro Gly Ile Trp Gln Leu Asp Cys Thr His Leu Glu Gly Lys Val Ile Leu Val Ala Val His Val Ala Ser Gly Tyr Ile Glu Ala Glu Val Ile Pro Ala Glu Thr Gly Gln Glu Thr Ala Tyr Phe Leu Leu Lys Leu Ala Gly Arg Trp Pro Val Lys Thr Ile His Thr Asp Asn Gly Ser Asn Phe Thr Gly Ala Thr Val 15 Arg Ala Ala Cys Trp Trp Ala Gly Ile Lys Gln Glu Phe Gly Ile Pro Tyr Asn Pro Gln Ser Gln Gly Val Val Glu Ser Met Asn Lys Glu Leu Lys Lys Ile Ile Gly Gln Val Arg Asp Gln Ala Glu His Leu Lys Thr Ala Val Gln Met Ala Val Phe Ile His Asn Phe Lys Arg Lys Gly Gly 20 Ile Gly Gly Tyr Ser Ala Gly Glu Arg Ile Val Asp Ile Ile Ala Thr Asp Ile Gln Thr Lys Glu Leu Gln Lys Gln Ile Thr Lys Ile Gln Asn Phe Arg Val Tyr Tyr Arg Asp Ser Arg Asn Pro Leu Trp Lys Gly Pro Ala Lys Leu Leu Trp Lys Gly Glu Gly Ala Val Val Ile Gln Asp Asn Ser Asp Ile Lys Val Val Pro Arg Arg Lys Ala Lys Ile Ile Arg Asp 25 Tyr Gly Lys Gln Met Ala Gly Asp Asp Cys Val Ala Ser Arg Gln Asp Glu Asp (SEQ ID NO:2).

The present invention especially relates to an adenoviral vector vaccine which comprises a codon optimized HIV-1 DNA pol construct wherein, in addition to deletion of the portion of the wild type sequence encoding the protease activity, a combination of active site residue mutations are introduced which are deleterious to HIV-1 pol (RT-RH-IN) activity of the expressed protein. Therefore, the present invention preferably relates to an adenoviral HIV-1 DNA pol-based vaccine wherein the construct is devoid of DNA sequences encoding any PR activity, as well as containing a mutation(s) which at least partially, and preferably substantially, abolishes RT, RNase and/or IN activity. One type of HIV-1 pol mutant which is part and parcel of an adenoviral vector vaccine may include but is not limited to a mutated

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DNA molecule comprising at least one nucleotide substitution which results in a point mutation which effectively alters an active site within the RT, RNase and/or IN regions of the expressed protein, resulting in at least substantially decreased enzymatic activity for the RT, RNase H and/or IN functions of HIV-1 Pol. In a preferred embodiment of this portion of the invention, a HIV-1 DNA pol construct contains a mutation or mutations within the Pol coding region which effectively abolishes RT, RNase H and IN activity. An especially preferable HIV-1 DNA pol construct in a DNA molecule which contains at least one point mutation which alters the active site of the RT, RNase H and IN domains of Pol, such that each activity is at least substantially abolished. Such a HIV-1 Pol mutant will most likely comprise at least one point mutation in or around each catalytic domain responsible for RT, RNase H and IN activity, respectfully. To this end, an especially preferred HIV-1 DNA pol construct is exemplified herein and contains nine codon substitution mutations which results in an inactivated Pol protein (IA Pol: SEQ ID NO:4, Figure 17A-C) which has no PR, RT, RNase or IN activity, wherein three such point mutations reside within each of the RT, RNase and IN catalytic domains. Therefore, an especially preferred exemplification is an adenoviral vaccine which comprises, in an appropriate fashion, a DNA molecule which encodes IA-pol, which contains all nine mutations as shown below in Table 1. An additional preferred amino acid residue for substitution is Asp551, localized within the RNase domain of Pol. Any combination of the mutations disclosed herein may suitable and therefore may be utilized as an IA-Pol-based vaccine of the present invention. While addition and deletion mutations are contemplated and within the scope of the invention, the preferred mutation is a point mutation resulting in a substitution of the wild type amino acid with an alternative amino acid residue.

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	11	116	

	wt aa	aa residue	mutant aa	enzyme function
	Asp	112	Ala	RT
	Asp	187	Ala	RT
35	Asp	188	Ala	RT
	Asp .	445	Ala	. RNase H
	Glu	480	Ala	RNase H
	Asp	500	Ala	RNase H
	Asp	626	Ala	IN
	Asp	678	Ala	IN
	Glu	714	Ala	IN

It is preferred that point mutations be incorporated into the IApol mutant adenoviral vaccines of the present invention so as to lessen the possibility of altering epitopes in and around the active site(s) of HIV-1 Pol.

To this end, SEQ ID NO:3 discloses the nucleotide sequence which codes for a codon optimized pol in addition to the nine mutations shown in Table 1, disclosed as follows, and referred to herein as "IApol":

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AGATCTACCA TGGCCCCCAT CTCCCCCATT GAGACTGTGC CTGTGAAGCT GAAGCCTGGC ATGGATGGCC CCAAGGTGAA GCAGTGGCCC CTGACTGAGG AGAAGATCAA GGCCCTGGTG GAAATCTGCA CTGAGATGGA GAAGGAGGGC AAAATCTCCA AGATTGGCCC CGAGAACCCC TACAACACCC CTGTGTTTGC CATCAAGAAG AAGGACTCCA CCAAGTGGAG GAAGCTGGTG GACTTCAGGG AGCTGAACAA GAGGACCCAG GACTTCTGGG AGGTGCAGCT GGGCATCCCC CACCCCGCTG GCCTGAAGAA GAAGAAGTCT GTGACTGTGC TGGCTGTGGG GGATGCCTAC TTCTCTGTGC CCCTGGATGA GGACTTCAGG AAGTACACTG CCTTCACCAT CCCCTCCATC AACAATGAGA CCCCTGGCAT CAGGTACCAG TACAATGTGC TGCCCCAGGG CTGGAAGGGC TCCCCTGCCA TCTTCCAGTC CTCCATGACC AAGATCCTGG AGCCCTTCAG GAAGCAGAAC CCTGACATTG TGATCTACCA GTACATGGCT GCCCTGTATG TGGGCTCTGA CCTGGAGATT GGGCAGCACA GGACCAAGAT TGAGGAGCTG AGGCAGCACC TGCTGAGGTG GGGCCTGACC ACCCCTGACA AGAAGCACCA GAAGGAGCCC CCCTTCCTGT GGATGGGCTA TGAGCTGCAC CCCGACAAGT GGACTGTGCA GCCCATTGTG CTGCCTGAGA AGGACTCCTG GACTGTGAAT GACATCCAGA AGCTGGTGGG CAAGCTGAAC TGGGCCTCCC AAATCTACCC TGGCATCAAG GTGAGGCAGC TGTGCAAGCT GCTGAGGGGC ACCAAGGCCC TGACTGAGGT GATCCCCCTG ACTGAGGAGG CTGAGCTGGA GCTGGCTGAG AACAGGGAGA TCCTGAAGGA GCCTGTGCAT GGGGTGTACT ATGACCCCTC CAAGGACCTG ATTGCTGAGA TCCAGAAGCA GGGCCAGGGC CAGTGGACCT ACCAAATCTA CCAGGAGCCC TTCAAGAACC TGAAGACTGG CAAGTATGCC AGGATGAGGG GGGCCCACAC CAATGATGTG AAGCAGCTGA CTGAGGCTGT GCAGAAGATC ACCACTGAGT CCATTGTGAT CTGGGGCAAG ACCCCCAAGT TCAAGCTGCC CATCCAGAAG GAGACCTGGG AGACCTGGTG GACTGAGTAC TGGCAGGCCA CCTGGATCCC TGAGTGGGAG TTTGTGAACA CCCCCCCCT GGTGAAGCTG TGGTACCAGC TGGAGAAGGA GCCCATTGTG GGGGCTGAGA CCTTCTATGT GGCTGGGGCT GCCAACAGGG AGACCAAGCT GGGCAAGGCT GGCTATGTGA CCAACAGGGG CAGGCAGAAG GTGGTGACCC TGACTGACAC CACCAACCAG AAGACTGCCC TCCAGGCCAT CTACCTGGCC CTCCAGGACT CTGGCCTGGA GGTGAACATT GTGACTGCCT CCCAGTATGC CCTGGGCATC ATCCAGGCCC AGCCTGATCA GTCTGAGTCT GTGCCTGCCC ACAAGGGCAT TGGGGGCAAT GAGCAGGTGG ACAAGCTGGT GTCTGCTGGC ATCAGGAAGG TGCTGTTCCT GGATGGCATT GACAAGGCCC AGGATGAGCA TGAGAAGTAC CACTCCAACT GGAGGGCTAT GGCCTCTGAC TTCAACCTGC CCCCTGTGGT GGCTAAGGAG

ATTGTGGCCT CCTGTGACAA GTGCCAGCTG AAGGGGGAGG CCATGCATGG GCAGGTGGAC TGCTCCCCTG GCATCTGGCA GCTGGCCTGC ACCCACCTGG AGGGCAAGGT GATCCTGGTG GCTGTGCATG TGGCCTCCGG CTACATTGAG GCTGAGGTGA TCCCTGCTGA GACAGGCCAG GAGACTGCCT ACTTCCTGCT GAAGCTGGCT GGCAGGTGGC CTGTGAAGAC CATCCACACT GCCAATGGCT CCAACTTCAC TGGGGCCACA GTGAGGGCTG CCTGCTGGTG GGCTGGCATC AAGCAGGAGT TTGGCATCCC CTACAACCCC CAGTCCCAGG GGGTGGTGGC CTCCATGAAC AAGGAGCTGA AGAAGATCAT TGGGCAGGTG AGGGACCAGG CTGAGCACCT GAAGACAGCT GTGCAGATGG CTGTGTTCAT CCACAACTTC AAGAGGAAGG GGGCATCGG GGGCTACTCC GCTGGGGAGA GGATTGTGGA CATCATTGCC ACAGACATCC AGACCAAGGA GCTCCAGAAG CAGATCACCA AGATCCAGAA CTTCAGGGTG TACTACAGGG ACTCCAGGAA CCCCCTGTGG AAGGGCCCTG CCAAGCTGCT GTGGAAGGGG GAGGGGGCTG TGGTGATCCA GGACAACTCT GACATCAAGG TGGTGCCCAG GAGGAAGGCC AAGATCATCA GGGACTATGG CAAGCAGATG GCTGGGGATG ACTGTGTGGC CTCCAGGCAG GATGAGGACT AAAGCCCGGG CAGATCT (SEQ ID NO:3).

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In order to produce the IA-pol-based adenoviral vaccines of the present invention, inactivation of the enzymatic functions was achieved by replacing a total of nine active site residues from the enzyme subunits with alanine side-chains. As shown in Table 1, all residues that comprise the catalytic triad of the polymerase, namely Asp112, Asp187, and Asp188, were substituted with alanine (Ala) residues (Larder, et al., Nature 1987, 327: 716-717; Larder, et al., 1989, Proc. Natl. Acad. Sci. 1989, 86: 4803-4807). Three additional mutations were introduced at Asp445, Glu480 and Asp500 to abolish RNase H activity (Asp551 was left unchanged in this IA Pol construct), with each residue being substituted for an Ala residue, respectively (Davies, et al., 1991, Science 252:, 88-95; Schatz, et al., 1989, FEBS Lett. 257: 311-314; Mizrahi, et al., 1990, Nucl. Acids. Res. 18: pp. 5359-5353). HIV pol integrase function was abolished through three mutations at Asp626, Asp678 and Glu714. Again, each of these residues has been substituted with an Ala residue (Wiskerchen, et al., 1995, J. Virol. 69: 376-386; Leavitt, et al., 1993, J. Biol. Chem. 268: 2113-2119). Amino acid residue Pro3 of SEQ ID NO:4 marks the start of the RT gene. The complete amino acid sequence of IA-Pol is disclosed herein as SEQ ID NO:4 and 30 Figure 17A-C, as follows:

Met Ala Pro Ile Ser Pro Ile Glu Thr Val Pro Val Lys Leu Lys Pro Gly Met Asp Gly Pro Lys Val Lys Gln Trp Pro Leu Thr Glu Glu Lys Ile Lys Ala Leu Val Glu Ile Cys Thr Glu Met Glu Lys Glu Gly Lys Ile Ser Lys Ile Gly Pro Glu Asn Pro Tyr Asn Thr Pro Val Phe Ala Ile Lys Lys Lys Asp Ser Thr Lys Trp Arg Lys Leu Val Asp Phe Arg

Glu Leu Asn Lys Arg Thr Gln Asp Phe Trp Glu Val Gln Leu Gly Ile Pro His Pro Ala Gly Leu Lys Lys Lys Ser Val Thr Val Leu Ala Val Gly Asp Ala Tyr Phe Ser Val Pro Leu Asp Glu Asp Phe Arg Lys Tyr Thr Ala Phe Thr Ile Pro Ser Ile Asn Asn Glu Thr Pro Gly Ile Arg Tyr Gln Tyr Asn Val Leu Pro Gln Gly Trp Lys Gly Ser Pro Ala Ile Phe Gln Ser Ser Met Thr Lys Ile Leu Glu Pro Phe Arg Lys Gln Asn Pro Asp Ile Val Ile Tyr Gln Tyr Met Ala Ala Leu Tyr Val Gly Ser Asp Leu Glu Ile Gly Gln His Arg Thr Lys Ile Glu Glu Leu Arg Gln His Leu Leu Arg Trp Gly Leu Thr Thr Pro Asp Lys Lys His Gln Lys Glu Pro Pro Phe Leu Trp Met Gly Tyr Glu Leu His Pro Asp Lys 10 Trp Thr Val Gln Pro Ile Val Leu Pro Glu Lys Asp Ser Trp Thr Val Asn Asp Ile Gln Lys Leu Val Gly Lys Leu Asn Trp Ala Ser Gln Ile Tyr Pro Gly Ile Lys Val Arg Gln Leu Cys Lys Leu Leu Arg Gly Thr Lys Ala Leu Thr Glu Val Ile Pro Leu Thr Glu Glu Ala Glu Leu Glu Leu Ala Glu Asn Arg Glu Ile Leu Lys Glu Pro Val His Gly Val Tyr 15 Tyr Asp Pro Ser Lys Asp Leu Ile Ala Glu Ile Gln Lys Gln Gly Gln Gly Gln Trp Thr Tyr Gln Ile Tyr Gln Glu Pro Phe Lys Asn Leu Lys Thr Gly Lys Tyr Ala Arg Met Arg Gly Ala His Thr Asn Asp Val Lys Gln Leu Thr Glu Ala Val Gln Lys Ile Thr Thr Glu Ser Ile Val Ile Trp Gly Lys Thr Pro Lys Phe Lys Leu Pro Ile Gln Lys Glu Thr Trp 20 Glu Thr Trp Trp Thr Glu Tyr Trp Gln Ala Thr Trp Ile Pro Glu Trp Glu Phe Val Asn Thr Pro Pro Leu Val Lys Leu Trp Tyr Gln Leu Glu Lys Glu Pro Ile Val Gly Ala Glu Thr Phe Tyr Val Ala Gly Ala Ala Asn Arg Glu Thr Lys Leu Gly Lys Ala Gly Tyr Val Thr Asn Arg Gly 25 Arg Gln Lys Val Val Thr Leu Thr Asp Thr Thr Asn Gln Lys Thr Ala Leu Gln Ala Ile Tyr Leu Ala Leu Gln Asp Ser Gly Leu Glu Val Asn Ile Val Thr Ala Ser Gln Tyr Ala Leu Gly Ile Ile Gln Ala Gln Pro Asp Gln Ser Glu Ser Glu Leu Val Asn Gln Ile Ile Glu Gln Leu Ile Lys Lys Glu Lys Val Tyr Leu Ala Trp Val Pro Ala His Lys Gly Ile 30 Gly Gly Asn Glu Gln Val Asp Lys Leu Val Ser Ala Gly Ile Arg Lys . Val Leu Phe Leu Asp Gly Ile Asp Lys Ala Gln Asp Glu His Glu Lys Tyr His Ser Asn Trp Arg Ala Met Ala Ser Asp Phe Asn Leu Pro Pro Val Val Ala Lys Glu Ile Val Ala Ser Cys Asp Lys Cys Gln Leu Lys Gly Glu Ala Met His Gly Gln Val Asp Cys Ser Pro Gly Ile Trp Gln Leu Ala Cys Thr His Leu Glu Gly Lys Val Ile Leu Val Ala Val His 35 Val Ala Ser Gly Tyr Ile Glu Ala Glu Val Ile Pro Ala Glu Thr Gly

Gln Glu Thr Ala Tyr Phe Leu Leu Lys Leu Ala Gly Arg Trp Pro Val Lys Thr Ile His Thr Ala Asn Gly Ser Asn Phe Thr Gly Ala Thr Val Arg Ala Ala Cys Trp Trp Ala Gly Ile Lys Gln Glu Phe Gly Ile Pro Tyr Asn Pro Gln Ser Gln Gly Val Val Ala Ser Met Asn Lys Glu Leu Lys Lys Ile Ile Gly Gln Val Arg Asp Gln Ala Glu His Leu Lys Thr Ala Val Gln Met Ala Val Phe Ile His Asn Phe Lys Arg Lys Gly Gly Ile Gly Gly Tyr Ser Ala Gly Glu Arg Ile Val Asp Ile Ile Ala Thr Asp Ile Gln Thr Lys Glu Leu Gln Lys Gln Ile Thr Lys Ile Gln Asn Phe Arg Val Tyr Tyr Arg Asp Ser Arg Asn Pro Leu Trp Lys Gly Pro Ala Lys Leu Leu Trp Lys Gly Glu Gly Ala Val Val Ile Gln Asp Asn Ser Asp Ile Lys Val Val Pro Arg Arg Lys Ala Lys Ile Ile Arg Asp Tyr Gly Lys Gln Met Ala Gly Asp Asp Cys Val Ala Ser Arg Gln Asp Glu Asp Glu Asp (SEQ ID NO:4).

As noted above, it will be understood that any combination of the mutations disclosed above may be suitable and therefore be utilized as an IA-pol-based adenoviral HIV vaccine of the present invention, either when administered alone or in a combined modality regime and/or a prime-boost regimen. For example, it may be possible to mutate only 2 of the 3 residues within the respective reverse transcriptase, RNase-H, and integrase coding regions while still abolishing these enzymatic activities. However, the IA-pol construct described above and disclosed as SEQ ID NO:3, as well as the expressed protein (SEQ ID NO:4;) is preferred. It is also preferred that at least one mutation be present in each of the three catalytic domains.

Another aspect of this portion of the invention are codon optimized HIV-1 Pol-based vaccine constructions which comprise a eukaryotic trafficking signal peptide such as from tPA (tissue-type plasminogen activator) or by a leader peptide such as is found in highly expressed mammalian proteins such as immunoglobulin leader peptides. Any functional leader peptide may be tested for efficacy. However, a preferred embodiment of the present invention, as with HIV-1 Nef constructs shown herein, is to provide for a HIV-1 Pol mutant adenoviral vaccine construction wherein the pol coding region or a portion thereof is operatively linked to a leader peptide, preferably a leader peptide from human tPA. In other words, a codon optimized HIV-1 Pol mutant such as IA-Pol (SEQ ID NO:4) may also comprise a leader peptide at the amino terminal portion of the protein, which may effect cellular trafficking and hence, immunogenicity of the expressed protein within the host cell. As noted in Figure 16A-B, a DNA vector which may be utilized to practice the present invention may be modified by known recombinant DNA methodology to contain a leader signal

peptide of interest, such that downstream cloning of the modified HIV-1 protein of interest results in a nucleotide sequence which encodes a modified HIV-1 tPA/Pol protein. In the alternative, as noted above, insertion of a nucleotide sequence which encodes a leader peptide may be inserted into a DNA vector housing the open reading frame for the Pol protein of interest. Regardless of the cloning strategy, the end result is a polynucleotide vaccine which comprises vector components for effective gene expression in conjunction with nucleotide sequences which encode a modified HIV-1 Pol protein of interest, including but not limited to a HIV-1 Pol protein which contains a leader peptide. The amino acid sequence of the human tPA leader utilized herein is as follows: MDAMKRGLCCVLLLCGAVFVSPSEISS (SEQ ID NO:17). Therefore, another aspect of the present invention is to generate HIV-1 Pol-based vaccine constructions which comprise a eukaryotic trafficking signal peptide such as from tPA. To this end, the present invention relates to a DNA molecule which encodes a codon optimized wt-pol DNA construct wherein the protease (PR) activity is deleted and a human tPA leader sequence is fused to the 5' end of the coding region. A DNA molecule which encodes this protein is disclosed herein as SEQ ID NO:5, the open reading frame disclosed herein as SEQ ID NO:6.

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To this end, the present invention relates to a DNA molecule which encodes a codon optimized wt-pol DNA construct wherein the protease (PR) activity is deleted and a human tPA leader sequence is fused to the 5' end of the coding region (herein, "tPA-wt-pol"). A DNA molecule which encodes this protein is disclosed herein as SEQ ID NO:5, the open reading frame being contained from an initiating Met residue at nucleotides 8-10 to a termination codon from nucleotides 2633-2635. SEQ ID NO:5 is as follows:

25 GATCACCATG GATGCAATGA AGAGAGGGCT CTGCTGTGT CTGCTGTGT GTGGAGCAGT
CTTCGTTTCG CCCAGCGAGA TCTCCGCCCC CATCTCCCC ATTGAGACTG TGCCTGTGAA
GCTGAAGCCT GGCATGGATG GCCCCAAGGT GAAGCAGTGG CCCCTGACTG AGGAGAAGAT
CAAGGCCCTG GTGGAAATCT GCACTGAGAT GGAGAAGGAG GGCAAAAATCT CCAAGATTGG
CCCCGAGAAC CCCTACAACA CCCCTGTGTT TGCCATCAAG AAGAAGGACT CCACCAAGTG
GCTGGGCATC CCCCACCCCG CTGGCCTGAA CAAGAGGACC CAGGACTTCT GGGAGGTGCA
GCGGGGATGCC TACTTCTCTG TGCCCCTGGA TGAGGACTTC AGGAAGTACA CTGCCTTCAC
CATCCCCTCC ATCAACAATG AGACCCCTGG CATCAGGTAC CAGTACAATG TGCTGCCCCA
GGGCTGGAAG GGCTCCCCTG CCATCTTCCA GTCCTCCATG ACCAAGATCC TGGAGCCCTT

35 CAGGAAGCAG AACCCTGACA TTGTGATCTA CCAGTACATG GATGACCTGT ATGTGGGCTC
TGACCTGGAG ATTGGGCAGC ACAGGACCAA GATTGAGGAG CTGAGGCAGC ACCTGCTGAG

GTGGGGCCTG ACCACCCCTG ACAAGAAGCA CCAGAAGGAG CCCCCCTTCC TGTGGATGGG CTATGAGCTG CACCCCGACA AGTGGACTGT GCAGCCCATT GTGCTGCCTG AGAAGGACTC CTGGACTGTG AATGACATCC AGAAGCTGGT GGGCAAGCTG AACTGGGCCT CCCAAATCTA CCCTGGCATC AAGGTGAGGC AGCTGTGCAA GCTGCTGAGG GGCACCAAGG CCCTGACTGA GGTGATCCCC CTGACTGAGG AGGCTGAGCT GGAGCTGGCT GAGAACAGGG AGATCCTGAA GGAGCCTGTG CATGGGGTGT ACTATGACCC CTCCAAGGAC CTGATTGCTG AGATCCAGAA GCAGGGCCAG GGCCAGTGGA CCTACCAAAT CTACCAGGAG CCCTTCAAGA ACCTGAAGAC TGGCAAGTAT GCCAGGATGA GGGGGGCCCA CACCAATGAT GTGAAGCAGC TGACTGAGGC TGTGCAGAAG ATCACCACTG AGTCCATTGT GATCTGGGGC AAGACCCCCA AGTTCAAGCT 10 GCCCATCCAG AAGGAGACCT GGGAGACCTG GTGGACTGAG TACTGGCAGG CCACCTGGAT CCCTGAGTGG GAGTTTGTGA ACACCCCCC CCTGGTGAAG CTGTGGTACC AGCTGGAGAA GGAGCCCATT GTGGGGGCTG AGACCTTCTA TGTGGATGGG GCTGCCAACA GGGAGACCAA GCTGGGCAAG GCTGGCTATG TGACCAACAG GGGCAGGCAG AAGGTGGTGA CCCTGACTGA CACCACCAAC CAGAAGACTG AGCTCCAGGC CATCTACCTG GCCCTCCAGG ACTCTGGCCT 15 GGAGGTGAAC ATTGTGACTG ACTCCCAGTA TGCCCTGGGC ATCATCCAGG CCCAGCCTGA TCAGTCTGAG TCTGAGCTGG TGAACCAGAT CATTGAGCAG CTGATCAAGA AGGAGAAGGT GTACCTGGCC TGGGTGCCTG CCCACAAGGG CATTGGGGGC AATGAGCAGG TGGACAAGCT GGTGTCTGCT GGCATCAGGA AGGTGCTGTT CCTGGATGGC ATTGACAAGG CCCAGGATGA GCATGAGAAG TACCACTCCA ACTGGAGGGC TATGGCCTCT GACTTCAACC TGCCCCCTGT 20 GGTGGCTAAG GAGATTGTGG CCTCCTGTGA CAAGTGCCAG CTGAAGGGGG AGGCCATGCA TGGGCAGGTG GACTGCTCCC CTGGCATCTG GCAGCTGGAC TGCACCCACC TGGAGGGCAA GGTGATCCTG GTGGCTGTGC ATGTGGCCTC CGGCTACATT GAGGCTGAGG TGATCCCTGC TGAGACAGGC CAGGAGACTG CCTACTTCCT GCTGAAGCTG GCTGGCAGGT GGCCTGTGAA GACCATCCAC ACTGACAATG GCTCCAACTT CACTGGGGCC ACAGTGAGGG CTGCCTGCTG 25 GTGGGCTGGC ATCAAGCAGG AGTTTGGCAT CCCCTACAAC CCCCAGTCCC AGGGGGTGGT GGAGTCCATG AACAAGGAGC TGAAGAAGAT CATTGGGCAG GTGAGGGACC AGGCTGAGCA CCTGAAGACA GCTGTGCAGA TGGCTGTGTT CATCCACAAC TTCAAGAGGA AGGGGGGCAT CGGGGGCTAC TCCGCTGGGG AGAGGATTGT GGACATCATT GCCACAGACA TCCAGACCAA GGAGCTCCAG AAGCAGATCA CCAAGATCCA GAACTTCAGG GTGTACTACA GGGACTCCAG 30 GAACCCCTG TGGAAGGGCC CTGCCAAGCT GCTGTGGAAG GGGGAGGGGG CTGTGGTGAT CCAGGACAAC TCTGACATCA AGGTGGTGCC CAGGAGGAAG GCCAAGATCA TCAGGGACTA TGGCAAGCAG ATGGCTGGGG ATGACTGTGT GGCCTCCAGG CAGGATGAGG ACTAAAGCCC GGGCAGATCT (SEQ ID NO:5).

The open reading frame of the wild type tPA-pol construct disclosed as SEQ ID NO:5 contains 875 amino acids, disclosed herein as SEQ ID NO:6, as follows:

Met Asp Ala Met Lys Arg Gly Leu Cys Cys Val Leu Leu Cys Gly

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Ala Val Phe Val Ser Pro Ser Glu Ile Ser Ala Pro Ile Ser Pro Ile Glu Thr Val Pro Val Lys Leu Lys Pro Gly Met Asp Gly Pro Lys Val Lys Gln Trp Pro Leu Thr Glu Glu Lys Ile Lys Ala Leu Val Glu Ile Cys Thr Glu Met Glu Lys Glu Gly Lys Ile Ser Lys Ile Gly Pro Glu 5 Asn Pro Tyr Asn Thr Pro Val Phe Ala Ile Lys Lys Lys Asp Ser Thr Lys Trp Arg Lys Leu Val Asp Phe Arg Glu Leu Asn Lys Arg Thr Gln Asp Phe Trp Glu Val Gln Leu Gly Ile Pro His Pro Ala Gly Leu Lys Lys Lys Lys Ser Val Thr Val Leu Asp Val Gly Asp Ala Tyr Phe Ser Val Pro Leu Asp Glu Asp Phe Arg Lys Tyr Thr Ala Phe Thr Ile Pro 10 Ser Ile Asn Asn Glu Thr Pro Gly Ile Arg Tyr Gln Tyr Asn Val Leu Pro Gln Gly Trp Lys Gly Ser Pro Ala Ile Phe Gln Ser Ser Met Thr Lys Ile Leu Glu Pro Phe Arg Lys Gln Asn Pro Asp Ile Val Ile Tyr Gln Tyr Met Asp Asp Leu Tyr Val Gly Ser Asp Leu Glu Ile Gly Gln His Arg Thr Lys Ile Glu Glu Leu Arg Gln His Leu Leu Arg Trp Gly 15 Leu Thr Thr Pro Asp Lys Lys His Gln Lys Glu Pro Pro Phe Leu Trp Met Gly Tyr Glu Leu His Pro Asp Lys Trp Thr Val Gln Pro Ile Val Leu Pro Glu Lys Asp Ser Trp Thr Val Asn Asp Ile Gln Lys Leu Val Gly Lys Leu Asn Trp Ala Ser Gln Ile Tyr Pro Gly Ile Lys Val Arg Gln Leu Cys Lys Leu Leu Arg Gly Thr Lys Ala Leu Thr Glu Val Ile 20 Pro Leu Thr Glu Glu Ala Glu Leu Glu Leu Ala Glu Asn Arg Glu Ile Leu Lys Glu Pro Val His Gly Val Tyr Tyr Asp Pro Ser Lys Asp Leu Ile Ala Glu Ile Gln Lys Gln Gly Gln Gly Gln Trp Thr Tyr Gln Ile Tyr Gln Glu Pro Phe Lys Asn Leu Lys Thr Gly Lys Tyr Ala Arg Met Arg Gly Ala His Thr Asn Asp Val Lys Gln Leu Thr Glu Ala Val Gln 25 Lys Ile Thr Thr Glu Ser Ile Val Ile Trp Gly Lys Thr Pro Lys Phe Lys Leu Pro Ile Gln Lys Glu Thr Trp Glu Thr Trp Trp Thr Glu Tyr Trp Gln Ala Thr Trp Ile Pro Glu Trp Glu Phe Val Asn Thr Pro Pro Leu Val Lys Leu Trp Tyr Gln Leu Glu Lys Glu Pro Ile Val Gly Ala Glu Thr Phe Tyr Val Asp Gly Ala Ala Asn Arg Glu Thr Lys Leu Gly 30 Lys Ala Gly Tyr Val Thr Asn Arg Gly Arg Gln Lys Val Val Thr Leu Thr Asp Thr Thr Asp Gln Lys Thr.Glu Leu Gln Ala Ile Tyr Leu Ala Leu Gln Asp Ser Gly Leu Glu Val Asn Ile Val Thr Asp Ser Gln Tyr Ala Leu Gly Ile Ile Gln Ala Gln Pro Asp Gln Ser Glu Ser Glu Leu Val Asn Gln Ile Ile Glu Gln Leu Ile Lys Lys Glu Lys Val Tyr Leu 35 Ala Trp Val Pro Ala His Lys Gly Ile Gly Gly Asn Glu Gln Val Asp Lys Leu Val Ser Ala Gly Ile Arg Lys Val Leu Phe Leu Asp Gly Ile

Asp Lys Ala Gln Asp Glu His Glu Lys Tyr His Ser Asn Trp Arg Ala Met Ala Ser Asp Phe Asn Leu Pro Pro Val Val Ala Lys Glu Ile Val Ala Ser Cys Asp Lys Cys Gln Leu Lys Gly Glu Ala Met His Gly Gln Val Asp Cys Ser Pro Gly Ile Trp Gln Leu Asp Cys Thr His Leu Glu Gly Lys Val Ile Leu Val Ala Val His Val Ala Ser Gly Tyr Ile Glu Ala Glu Val Ile Pro Ala Glu Thr Gly Gln Glu Thr Ala Tyr Phe Leu Leu Lys Leu Ala Gly Arg Trp Pro Val Lys Thr Ile His Thr Asp Asn Gly Ser Asn Phe Thr Gly Ala Thr Val Arg Ala Ala Cys Trp Trp Ala Gly Ile Lys Gln Glu Phe Gly Ile Pro Tyr Asn Pro Gln Ser Gln Gly Val Val Glu Ser Met Asn Lys Glu Leu Lys Lys Ile Ile Gly Gln Val Arg Asp Gln Ala Glu His Leu Lys Thr Ala Val Gln Met Ala Val Phe Ile His Asn Phe Lys Arg Lys Gly Gly Ile Gly Gly Tyr Ser Ala Gly Glu Arg Ile Val Asp Ile Ile Ala Thr Asp Ile Gln Thr Lys Glu Leu Gln Lys Gln Ile Thr Lys Ile Gln Asn Phe Arg Val Tyr Tyr Arg Asp Ser Arg Asn Pro Leu Trp Lys Gly Pro Ala Lys Leu Leu Trp Lys Gly Glu Gly Ala Val Val Ile Gln Asp Asn Ser Asp Ile Lys Val Val Pro Arg Arg Lys Ala Lys Ile Ile Arg Asp Tyr Gly Lys Gln Met Ala Gly Asp Asp Cys Val Ala Ser Arg Gln Asp Glu Asp (SEQ ID NO:6).

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The present invention also relates to a codon optimized HIV-1 Pol mutant contained within a recombinant adenoviral vector such as IA-Pol (SEQ ID NO:4) which comprises a leader peptide at the amino terminal portion of the protein, which may effect cellular trafficking and hence, immunogenicity of the expressed protein within the host cell. Any such adenoviral-based HIV-1 DNA pol mutant disclosed in the above paragraphs is suitable for fusion downstream of a leader peptide, such as a leader peptide including but not limited to the human tPA leader sequence. Therefore, any such leader peptide-based HIV-1 pol mutant construct may include but is not limited to a mutated DNA molecule which effectively alters the catalytic activity of the RT, RNase and/or IN region of the expressed protein, resulting in at least substantially decreased enzymatic activity one or more of the RT, RNase H and/or IN functions of HIV-1 Pol. In a preferred embodiment of this portion of the invention, a leader peptide/HIV-1 DNA pol construct contains a mutation or mutations within the Pol coding region which effectively abolishes RT, RNase H and IN activity. An especially preferable HIV-1 DNA pol construct is a DNA molecule which contains at least one point mutation which alters the active site and catalytic activity within the RT, RNase H and IN domains of Pol, such that each activity is at least substantially abolished, and preferably totally abolished. Such a HIV-1 Pol mutant will most likely

comprise at least one point mutation in or around each catalytic domain responsible for RT, RNase H and IN activity, respectfully. An especially preferred embodiment of this portion of the invention relates to a human tPA leader fused to the IA-Pol protein comprising the nine mutations shown in Table 1. The DNA molecule is disclosed herein as SEQ ID NO:7 and the expressed tPA-IA Pol protein comprises a fusion junction as shown in Figure 18. The complete amino acid sequence of the expressed protein is set forth in SEQ ID NO:8. To this end, SEQ ID NO:7 discloses the nucleotide sequence which codes for a human tPA leader fused to the IA Pol protein comprising the nine mutations shown in Table 1 (herein, "tPA-opt-IApol"). The open reading frame begins with the initiating Met (nucleotides 8-10) and terminates with a "TAA" codon at nucleotides 2633-2635. The nucleotide sequence encoding tPA-IAPol is also disclosed as follows: GATCACCATG GATGCAATGA AGAGAGGGCT CTGCTGTGTG CTGCTGCTGT GTGGAGCAGT CTTCGTTTCG CCCAGCGAGA TCTCCGCCCC CATCTCCCCC ATTGAGACTG TGCCTGTGAA GCTGAAGCCT GGCATGGATG GCCCCAAGGT GAAGCAGTGG CCCCTGACTG AGGAGAAGAT CAAGGCCCTG GTGGAAATCT GCACTGAGAT GGAGAAGGAG GGCAAAATCT CCAAGATTGG CCCCGAGAAC CCCTACAACA CCCCTGTGTT TGCCATCAAG AAGAAGGACT CCACCAAGTG GAGGAAGCTG GTGGACTTCA GGGAGCTGAA CAAGAGGACC CAGGACTTCT GGGAGGTGCA GCTGGGCATC CCCCACCCCG CTGGCCTGAA GAAGAAGAAG TCTGTGACTG TGCTGGCTGT GGGGGATGCC TACTTCTCTG TGCCCCTGGA TGAGGACTTC AGGAAGTACA CTGCCTTCAC CATCCCCTCC ATCAACAATG AGACCCCTGG CATCAGGTAC CAGTACAATG TGCTGCCCCA GGGCTGGAAG GGCTCCCCTG CCATCTTCCA GTCCTCCATG ACCAAGATCC TGGAGCCCTT CAGGAAGCAG AACCCTGACA TTGTGATCTA CCAGTACATG GCTGCCCTGT ATGTGGGCTC TGACCTGGAG ATTGGGCAGC ACAGGACCAA GATTGAGGAG CTGAGGCAGC ACCTGCTGAG GTGGGGCCTG ACCACCCCTG ACAAGAAGCA CCAGAAGGAG CCCCCCTTCC TGTGGATGGG CTATGAGCTG CACCCCGACA AGTGGACTGT GCAGCCCATT GTGCTGCCTG AGAAGGACTC CTGGACTGTG AATGACATCC AGAAGCTGGT GGGCAAGCTG AACTGGGCCT CCCAAATCTA CCCTGGCATC AAGGTGAGGC AGCTGTGCAA GCTGCTGAGG GGCACCAAGG CCCTGACTGA GGTGATCCCC CTGACTGAGG AGGCTGAGCT GGAGCTGGCT GAGAACAGGG AGATCCTGAA GGAGCCTGTG CATGGGGTGT ACTATGACCC CTCCAAGGAC CTGATTGCTG AGATCCAGAA -GCAGGGCCAG GGCCAGTGGA CCTACCAAAT CTACCAGGAG CCCTTCAAGA ACCTGAAGAC TGGCAAGTAT GCCAGGATGA GGGGGCCCA CACCAATGAT GTGAAGCAGC TGACTGAGGC TGTGCAGAAG ATCACCACTG AGTCCATTGT GATCTGGGGC AAGACCCCCA AGTTCAAGCT GCCCATCCAG AAGGAGACCT GGGAGACCTG GTGGACTGAG TACTGGCAGG CCACCTGGAT CCCTGAGTGG GAGTTTGTGA ACACCCCCC CCTGGTGAAG CTGTGGTACC AGCTGGAGAA GGAGCCCATT GTGGGGGCTG AGACCTTCTA TGTGGCTGGG GCTGCCAACA GGGAGACCAA

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GCTGGGCAAG GCTGGCTATG TGACCAACAG GGGCAGGCAG AAGGTGGTGA CCCTGACTGA CACCACCAAC CAGAAGACTG CCCTCCAGGC CATCTACCTG GCCCTCCAGG ACTCTGGCCT GGAGGTGAAC ATTGTGACTG CCTCCCAGTA TGCCCTGGGC ATCATCCAGG CCCAGCCTGA TCAGTCTGAG TCTGAGCTGG TGAACCAGAT CATTGAGCAG CTGATCAAGA AGGAGAAGGT GTACCTGGCC TGGGTGCCTG CCCACAAGGG CATTGGGGGC AATGAGCAGG TGGACAAGCT GGTGTCTGCT GGCATCAGGA AGGTGCTGTT CCTGGATGGC ATTGACAAGG CCCAGGATGA GCATGAGAAG TACCACTCCA ACTGGAGGGC TATGGCCTCT GACTTCAACC TGCCCCCTGT GGTGGCTAAG GAGATTGTGG CCTCCTGTGA CAAGTGCCAG CTGAAGGGGG AGGCCATGCA TGGGCAGGTG GACTGCTCCC CTGGCATCTG GCAGCTGGCC TGCACCCACC TGGAGGGCAA GGTGATCCTG GTGGCTGTGC ATGTGGCCTC CGGCTACATT GAGGCTGAGG TGATCCCTGC TGAGACAGGC CAGGAGACTG CCTACTTCCT GCTGAAGCTG GCTGGCAGGT GGCCTGTGAA GACCATCCAC ACTGCCAATG GCTCCAACTT CACTGGGGCC ACAGTGAGGG CTGCCTGCTG GTGGGCTGGC ATCAAGCAGG AGTTTGGCAT CCCCTACAAC CCCCAGTCCC AGGGGGTGGT GGCCTCCATG AACAAGGAGC TGAAGAAGAT CATTGGGCAG GTGAGGGACC AGGCTGAGCA CCTGAAGACA GCTGTGCAGA TGGCTGTGTT CATCCACAAC TTCAAGAGGA AGGGGGGCAT CGGGGGCTAC TCCGCTGGGG AGAGGATTGT GGACATCATT GCCACAGACA TCCAGACCAA GGAGCTCCAG AAGCAGATCA CCAAGATCCA GAACTTCAGG GTGTACTACA GGGACTCCAG GAACCCCCTG TGGAAGGGCC CTGCCAAGCT GCTGTGGAAG GGGGAGGGGG CTGTGGTGAT CCAGGACAAC TCTGACATCA AGGTGGTGCC CAGGAGGAAG GCCAAGATCA TCAGGGACTA TGGCAAGCAG ATGGCTGGGG ATGACTGTGT GGCCTCCAGG CAGGATGAGG ACTAAAGCCC 20 GGGCAGATCT (SEQ ID NO:7).

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The open reading frame of the tPA-IA-pol construct disclosed as SEQ ID NO:7 contains 875 amino acids, disclosed herein as tPA-IA-Pol and SEQ ID NO:8, as follows:

Met Asp Ala Met Lys Arg Gly Leu Cys Cys Val Leu Leu Cys Gly 25 Ala Val Phe Val Ser Pro Ser Glu Ile Ser Ala Pro Ile Ser Pro Ile Glu Thr Val Pro Val Lys Leu Lys Pro Gly Met Asp Gly Pro Lys Val Lys Gln Trp Pro Leu Thr Glu Glu Lys Ile Lys Ala Leu Val Glu Ile Cys Thr Glu Met Glu Lys Glu Gly Lys Ile Ser Lys Ile Gly Pro Glu Asn Pro Tyr Asn Thr Pro Val Phe Ala Ile Lys Lys Asp Ser Thr 30 Lys Trp Arg Lys Leu Val Asp Phe Arg Glu Leu Asn Lys Arg Thr Gln Asp Phe Trp Glu Val Gln Leu Gly Ile Pro His Pro Ala Gly Leu Lys Lys Lys Lys Ser Val Thr Val Leu Ala Val Gly Asp Ala Tyr Phe Ser Val Pro Leu Asp Glu Asp Phe Arg Lys Tyr Thr Ala Phe Thr Ile Pro Ser Ile Asn Asn Glu Thr Pro Gly Ile Arg Tyr Gln Tyr Asn Val Leu 35 Pro Gln Gly Trp Lys Gly Ser Pro Ala Ile Phe Gln Ser Ser Met Thr

Lys Ile Leu Glu Pro Phe Arg Lys Gln Asn Pro Asp Ile Val Ile Tyr Gln Tyr Met Ala Ala Leu Tyr Val Gly Ser Asp Leu Glu Ile Gly Gln His Arg Thr Lys Ile Glu Glu Leu Arg Gln His Leu Leu Arg Trp Gly Leu Thr Thr Pro Asp Lys Lys His Gln Lys Glu Pro Pro Phe Leu Trp Met Gly Tyr Glu Leu His Pro Asp Lys Trp Thr Val Gln Pro Ile Val Leu Pro Glu Lys Asp Ser Trp Thr Val Asn Asp Ile Gln Lys Leu Val Gly Lys Leu Asn Trp Ala Ser Gln Ile Tyr Pro Gly Ile Lys Val Arg Gln Leu Cys Lys Leu Leu Arg Gly Thr Lys Ala Leu Thr Glu Val Ile Pro Leu Thr Glu Glu Ala Glu Leu Glu Leu Ala Glu Asn Arg Glu Ile Leu Lys Glu Pro Val His Gly Val Tyr Tyr Asp Pro Ser Lys Asp Leu 10 Ile Ala Glu Ile Gln Lys Gln Gly Gln Gly Gln Trp Thr Tyr Gln Ile Tyr Gln Glu Pro Phe Lys Asn Leu Lys Thr Gly Lys Tyr Ala Arg Met Arg Gly Ala His Thr Asn Asp Val Lys Gln Leu Thr Glu Ala Val Gln Lys Ile Thr Thr Glu Ser Ile Val Ile Trp Gly Lys Thr Pro Lys Phe Lys Leu Pro Ile Gln Lys Glu Thr Trp Glu Thr Trp Trp Thr Glu Tyr 15 Trp Gln Ala Thr Trp Ile Pro Glu Trp Glu Phe Val Asn Thr Pro Pro Leu Val Lys Leu Trp Tyr Gln Leu Glu Lys Glu Pro Ile Val Gly Ala Glu Thr Phe Tyr Val Ala Gly Ala Ala Asn Arg Glu Thr Lys Leu Gly Lys Ala Gly Tyr Val Thr Asn Arg Gly Arg Gln Lys Val Val Thr Leu Thr Asp Thr Thr Asn Gln Lys Thr Ala Leu Gln Ala Ile Tyr Leu Ala 20 Leu Gln Asp Ser Gly Leu Glu Val Asn Ile Val Thr Ala Ser Gln Tyr Ala Leu Gly Ile Ile Gln Ala Gln Pro Asp Gln Ser Glu Ser Glu Leu Val Asn Gln Ile Ile Glu Gln Leu Ile Lys Lys Glu Lys Val Tyr Leu Ala Trp Val Pro Ala His Lys Gly Ile Gly Gly Asn Glu Gln Val Asp Lys Leu Val Ser Ala Gly Ile Arg Lys Val Leu Phe Leu Asp Gly Ile 25 Asp Lys Ala Gln Asp Glu His Glu Lys Tyr His Ser Asn Trp Arg Ala Met Ala Ser Asp Phe Asn Leu Pro Pro Val Val Ala Lys Glu Ile Val Ala Ser Cys Asp Lys Cys Gln Leu Lys Gly Glu Ala Met His Gly Gln Val Asp Cys Ser Pro Gly Ile Trp Gln Leu Ala Cys Thr His Leu Glu 30 Gly Lys Val Ile Leu Val Ala Val His Val Ala Ser Gly Tyr Ile Glu Ala Glu Val Ile Pro Ala Glu Thr Gly Gln Glu Thr Ala Tyr Phe Leu Leu Lys Leu Ala Gly Arg Trp Pro Val Lys Thr Ile His Thr Ala Asn Gly Ser Asn Phe Thr Gly Ala Thr Val Arg Ala Ala Cys Trp Trp Ala Gly Ile Lys Gln Glu Phe Gly Ile Pro Tyr Asn Pro Gln Ser Gln Gly Val Val Ala Ser Met Asn Lys Glu Leu Lys Lys Ile Ile Gly Gln Val 35 Arg Asp Gln Ala Glu His Leu Lys Thr Ala Val Gln Met Ala Val Phe

Ile His Asn Phe Lys Arg Lys Gly Gly Ile Gly Gly Tyr Ser Ala Gly Glu Arg Ile Val Asp Ile Ile Ala Thr Asp Ile Gln Thr Lys Glu Leu Gln Lys Gln Ile Thr Lys Ile Gln Asn Phe Arg Val Tyr Tyr Arg Asp Ser Arg Asn Pro Leu Trp Lys Gly Pro Ala Lys Leu Leu Trp Lys Gly Glu Gly Ala Val Val Ile Gln Asp Asn Ser Asp Ile Lys Val Val Pro Arg Arg Lys Ala Lys Ile Ile Arg Asp Tyr Gly Lys Gln Met Ala Gly Asp Asp Cys Val Ala Ser Arg Gln Asp Glu Asp (SEQ ID NO:8).

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EXAMPLE 18

CODON OPTIMIZED HIV-1 NEF AND CODON OPTIMIZED HIV-1 NEF MODIFICATIONS

Codon optimized version of HIV-1 Nef and HIV-1 Nef modifications are essentially as described in U.S. Application Serial No. 09/738,782, filed December 15, 2000 and PCT International Application PCT/US00/34162, also filed December 15, 2000, both documents which are hereby incorporated by reference. As disclosed within the above-mentioned documents, particular embodiments of codon optimized Nef and Nef modifications relate to a DNA molecule encoding HIV-1 Nef from the HIV-1 jfrl isolate wherein the codons are optimized for expression in a mammalian system such as a human. The DNA molecule which encodes this protein is disclosed herein as SEQ ID NO:9, while the expressed open reading frame is disclosed herein as SEQ ID NO:10. Another embodiment of Nef-based coding regions for use in the adenoviral vectors of the present invention comprise a codon optimized DNA molecule encoding a protein containing the human plasminogen activator (tpa) leader peptide fused with the NH2-terminus of the HIV-1 Nef polypeptide. The DNA molecule which encodes this protein is disclosed herein as SEQ ID NO:11, while the expressed open reading frame is disclosed herein as SEQ ID NO:12. Another modified Nef optimized coding region relates to a DNA molecule encoding optimized HIV-1 Nef wherein the open reading frame codes for modifications at the amino terminal myristylation site (Gly-2 to Ala-2) and substitution of the Leu-174-Leu-175 dileucine motif to Ala-174-Ala-175, herein described as opt nef (G2A, LLAA). The DNA molecule which encodes this protein is disclosed herein as SEQ ID NO:13, while the expressed open reading frame is disclosed herein as SEQ ID NO:14. An additional embodiment relates to a DNA molecule encoding optimized HIV-1 Nef wherein the amino terminal myristylation site and dileucine motif have been deleted, as well as comprising a tPA leader peptide. This DNA molecule, opt tpanef (LLAA), comprises an open reading frame which

encodes a Nef protein containing a tPA leader sequence fused to amino acid residue 6-216 of HIV-1 Nef (jfrl), wherein Leu-174 and Leu-175 are substituted with Ala-174 and Ala-175, herein referred to as opt tpanef (LLAA) is disclosed herein as SEQ ID NO:15, while the expressed open reading frame is disclosed herein as SEQ ID NO:16.

As disclosed in the above-identified documents (U.S. Application Serial No. 09/738,782 and PCT International Application PCT/US00/34162) and reiterated herein, the following nef-based nucleotide and amino acid sequences which comprise the respective open reading frame are as follows:

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The nucleotide sequence of the codon optimized version of HIV-1 jrfl nef gene is disclosed herein as SEQ ID NO:9, as shown herein:

GATCTGCCAC CATGGGCGGC AAGTGGTCCA AGAGGTCCGT GCCCGGCTGG TCCACCGTGA GGGAGAGGAT GAGGAGGGCC GAGCCCGCCG CCGACAGGGT GAGGAGGACC GAGCCCGCCG CCGTGGGCGT GGGCGCCGTG TCCAGGGACC TGGAGAAGCA CGGCGCCATC ACCTCCTCCA ACACCGCCGC CACCAACGCC GACTGCGCCT GGCTGGAGGC CCAGGAGGAC GAGGAGGTGG GCTTCCCCGT GAGGCCCCAG GTGCCCCTGA GGCCCATGAC CTACAAGGGC GCCGTGGACC TGTCCCACTT CCTGAAGGAG AAGGGCGGCC TGGAGGGCCT GATCCACTCC CAGAAGAGGC AGGACATCCT GGACCTGTGG GTGTACCACA CCCAGGGCTA CTTCCCCGAC TGGCAGAACT ACACCCCGG CCCCGGCATC AGGTTCCCCC TGACCTTCGG CTGGTGCTTC AAGCTGGTGC CCGTGGAGCC CGAGAAGGTG GAGGAGGCCA ACGAGGGCGA GAACAACTGC CTGCTGCACC CCATGTCCCA GCACGGCATC GAGGACCCCG AGAAGGAGGT GCTGGAGTGG AGGTTCGACT CCAAGCTGGC CTTCCACCAC GTGGCCAGGG AGCTGCACCC CGAGTACTAC AAGGACTGCT AAAGCCCGGG C (SEQ ID NO:9).

Preferred codon usage is as follows: Met (ATG), Gly (GGC), Lys (AAG), Trp (TGG), Ser (TCC), Arg (AGG), Val (GTG), Pro (CCC), Thr (ACC), Glu (GAG); Leu (CTG), His (CAC), Ile (ATC), Asn (AAC), Cys (TGC), Ala (GCC), Gln (CAG), Phe (TTC) and Tyr (TAC). For an additional discussion relating to mammalian (human) codon optimization, see WO 97/31115 (PCT/US97/02294), which is hereby incorporated by reference. See also Figure 19A-B for a comparion of wild type vs. codon optimized nucleotides comprising the open reading frame of HIV-Nef.

The open reading frame for SEQ ID NO:9 above comprises an initiating methionine residue at nucleotides 12-14 and a "TAA" stop codon from nucleotides 660-662. The open reading frame of SEQ ID NO:9 provides for a 216 amino acid HIV-1 Nef protein expressed through utilization of a codon optimized DNA vaccine vector. The 216 amino acid HIV-1 Nef (jfrl) protein is disclosed herein as SEQ ID 35 NO:10, and as follows:

Met Gly Gly Lys Trp Ser Lys Arg Ser Val Pro Gly Trp Ser Thr Val

Arg Glu Arg Met Arg Arg Ala Glu Pro Ala Ala Asp Arg Val Arg Arg Thr Glu Pro Ala Ala Val Gly Val Gly Ala Val Ser Arg Asp Leu Glu Lys His Gly Ala Ile Thr Ser Ser Asn Thr Ala Ala Thr Asn Ala Asp Cys Ala Trp Leu Glu Ala Gln Glu Asp Glu Glu Val Gly Phe Pro Val Arg Pro Gln Val Pro Leu Arg Pro Met Thr Tyr Lys Gly Ala Val Asp Leu Ser His Phe Leu Lys Glu Lys Gly Gly Leu Glu Gly Leu Ile His Ser Gln Lys Arg Gln Asp Ile Leu Asp Leu Trp Val Tyr His Thr Gln Gly Tyr Phe Pro Asp Trp Gln Asn Tyr Thr Pro Gly Pro Gly Ile Arg Phe Pro Leu Thr Phe Gly Trp Cys Phe Lys Leu Val Pro Val Glu Pro Glu Lys Val Glu Glu Ala Asn Glu Gly Glu Asn Asn Cys Leu Leu His Pro Met Ser Gln His Gly Ile Glu Asp Pro Glu Lys Glu Val Leu Glu Trp Arg Phe Asp Ser Lys Leu Ala Phe His His Val Ala Arg Glu Leu His Pro Glu Tyr Tyr Lys Asp Cys (SEQ ID NO:10).

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HIV-1 Nef is a 216 amino acid cytosolic protein which associates with the inner surface of the host cell plasma membrane through myristylation of Gly-2 (Franchini et al., 1986, Virology 155: 593-599). While not all possible Nef functions have been elucidated, it has become clear that correct trafficking of Nef to the inner plasma membrane promotes viral replication by altering the host intracellular environment to facilitate the early phase of the HIV-1 life cycle and by increasing the infectivity of progeny viral particles. In one aspect of the invention regarding codon-optimized, protein-modified polypeptides, the nef-encoding region of the adenovirus vector of the present invention is modified to contain a nucleotide sequence which encodes a heterologous leader peptide such that the amino terminal region of the expressed protein will contain the leader peptide. The diversity of function that typifies eukaryotic cells depends upon the structural differentiation of their membrane boundaries. To generate and maintain these structures, proteins must be transported from their site of synthesis in the endoplasmic reticulum to predetermined destinations throughout the cell. This requires that the trafficking proteins display sorting signals that are recognized by the molecular machinery responsible for route selection located at the access points to the main trafficking pathways. Sorting decisions for most proteins need to be made only once as they traverse their biosynthetic pathways since their final destination, the cellular location at which they perform their function, becomes their permanent residence. Maintenance of intracellular integrity depends in part on the selective sorting and accurate transport of proteins to their correct destinations. Defined sequence motifs exist in proteins which can act as 'address labels'. A number of sorting signals have

been found associated with the cytoplasmic domains of membrane proteins. An effective induction of CTL responses often required sustained, high level endogenous expression of an antigen. As membrane-association via myristylation is an essential requirement for most of Nef's function, mutants lacking myristylation, by glycine-to-alanine change, change of the dileucine motif and/or by substitution with a tpa leader sequence as described herein, will be functionally defective, and therefore will have improved safety profile compared to wild-type Nef for use as an HIV-1 vaccine component.

In another embodiment of this portion of the invention, either the DNA vector or the HIV-1 nef nucleotide sequence is modified to include the human tissue-specific plasminogen activator (tPA) leader. As shown in Figure 16A-B, a DNA vector may be modified by known recombinant DNA methodology to contain a leader signal peptide of interest, such that downstream cloning of the modified HIV-1 protein of interest results in a nucleotide sequence which encodes a modified HIV-1 tPA/Nef protein. In the alternative, as noted above, insertion of a nucleotide sequence which encodes a leader peptide may be inserted into a DNA vector housing the open reading frame for the Nef protein of interest. Regardless of the cloning strategy, the end result is a polynucleotide vaccine which comprises vector components for effective gene expression in conjunction with nucleotide sequences which encode a modified HIV-1 Nef protein of interest, including but not limited to a HIV-1 Nef protein which contains a leader peptide. The amino acid sequence of the human tPA leader utilized herein is as follows: MDAMKRGLCCVLLLCGAVFVSPSEISS (SEQ ID NO:17).

It has been shown that myristylation of Gly-2 in conjunction with a dileucine motif in the carboxy region of the protein is essential for Nef-induced down regulation of CD4 (Aiken et al., 1994, Cell 76: 853-864) via endocytosis. It has also been shown that Nef expression promotes down regulation of MHCI (Schwartz et al., 1996, Nature Medicine 2(3): 338-342) via endocytosis. The present invention relates in part to DNA vaccines which encode modified Nef proteins altered in trafficking and/or functional properties. The modifications introduced into the adenoviral vector HIV vaccines of the present invention include but are not limited to additions, deletions or substitutions to the nef open reading frame which results in the expression of a modified Nef protein which includes an amino terminal leader peptide, modification or deletion of the amino terminal myristylation site, and modification or deletion of the dileucine motif within the Nef protein and which alter function within the infected host cell. Therefore, a central theme of the DNA molecules and recombinant adenoviral HIV vaccines of the present invention is (1)

host administration and intracellular delivery of a codon optimized nef-based adenoviral HIV vaccine; (2) expression of a modified Nef protein which is immunogenic in terms of eliciting both CTL and Th responses; and, (3) inhibiting or at least altering known early viral functions of Nef which have been shown to promote HIV-1 replication and load within an infected host. Therefore, the nef coding region may be altered, resulting in a DNA vaccine which expresses a modified Nef protein wherein the amino terminal Gly-2 myristylation residue is either deleted or modified to express alternate amino acid residues. Also, the nef coding region may be altered so as to result in a DNA vaccine which expresses a modified Nef protein wherein the dileucine motif is either deleted or modified to express alternate amino acid residues. In addition, the adenoviral vector HIV vaccines of the present invention also relate to an isolated DNA molecule, regardless of codon usage, which expresses a wild type or modified Nef protein as described herein, including but not limited to modified Nef proteins which comprise a deletion or substitution of Gly 2, a deletion or substitution of Leu 174 and Leu 175 and/or inclusion of a leader sequence.

Therefore, specific Nef-based constructs further include the following, as exemplification's and not limitations. For example, the present invention relates to an adenoviral vector vaccine which encodes modified forms of HIV-1, an open reading frame which encodes a Nef protein which comprises a tPA leader sequence fused to amino acid residue 6-216 of HIV-1 Nef (jfrl) is referred to herein as opt tpanef. The nucleotide sequence comprising the open reading frame of opt tpanef is disclosed herein as SEQ ID NO:11, as shown below:

CATGGATGCA ATGAAGAGA GGCTCTGCTG TGTGCTGCTG CTGTGTGGAG CAGTCTTCGT
TTCGCCCAGC GAGATCTCCT CCAAGAGGTC CGTGCCCGGC TGGTCCACCG TGAGGGAGAG
GATGAGGAGG GCCGAGCCCG CCGCCGACAG GGTGAGGAGG ACCGAGCCCG CCGCCGTGGG
CGTGGGCGCC GTGTCCAGGG ACCTGGAGAA GCACGGCGCC ATCACCTCCT CCAACACCGC
CGCCACCAAC GCCGACTGCG CCTGGCTGGA GGCCCAGGAG GACGAGGAGG TGGGCTTCCC
CGTGAGGCCC CAGGTGCCCC TGAGGCCCAT GACCTACAAG GGCGCCGTGG ACCTGTCCCA
CTTCCTGAAG GAGAAGGGCG GCCTGGAGGG CCTGATCCAC TCCCAGAAGA GGCAGGACAT
CCTGGACCTG TGGGTGTACC ACACCCAGGG CTACTTCCCC GACTGGCAGA ACTACACCCC
CGGCCCCGGC ATCAGGTTCC CCCTGACCTT CGGCTGTGC TTCAAGCTGG TGCCCGTGGA
GCCCGAGAAG GTGGAGGAGG CCAACGAGGG CGAGAACAAC TGCCTGCTGC ACCCCATGTC
CCAGCACGGC ATCGAGGACC CCGAGAAGGA GGTGCTGGAG TGGAGGTTCG ACCCCATGTC
GGCCTTCCAC CACGTGGCCA GGGAGCTGCA CCCCGAGTAC TACAAGGACT GCTAAAGCC
(SEQ ID NO:11).

The open reading frame for SEQ ID NO:11 comprises an initiating methionine

residue at nucleotides 2-4 and a "TAA" stop codon from nucleotides 713-715. The open reading frame of SEQ ID NO:3 provides for a 237 amino acid HIV-1 Nef protein which comprises a tPA leader sequence fused to amino acids 6-216 of HIV-1 Nef, including the dileucine motif at amino acid residues 174 and 175. This 237 amino acid tPA/Nef (jfrl) fusion protein is disclosed herein as SEQ ID NO:12, and is shown as follows:

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Met Asp Ala Met Lys Arg Gly Leu Cys Cys Val Leu Leu Cys Gly Ala Val Phe Val Ser Pro Ser Glu Ile Ser Ser Lys Arg Ser Val Pro Gly Trp Ser Thr Val Arg Glu Arg Met Arg Arg Ala Glu Pro Ala Ala Asp Arg Val Arg Arg Thr Glu Pro Ala Ala Val Gly Val Gly Ala Val Ser Arg Asp Leu Glu Lys His Gly Ala Ile Thr Ser Ser Asn Thr Ala Ala Thr Asn Ala Asp Cys Ala Trp Leu Glu Ala Gln Glu Asp Glu Glu Val Gly Phe Pro Val Arg Pro Gln Val Pro Leu Arg Pro Met Thr Tyr Lys Gly Ala Val Asp Leu Ser His Phe Leu Lys Glu Lys Gly Gly Leu Glu Gly Leu Ile His Ser Gln Lys Arg Gln Asp Ile Leu Asp Leu Trp Val Tyr His Thr Gln Gly Tyr Phe Pro Asp Trp Gln Asn Tyr Thr Pro Gly Pro Gly Ile Arg Phe Pro Leu Thr Phe Gly Trp Cys Phe Lys Leu Val Pro Val Glu Pro Glu Lys Val Glu Glu Ala Asn Glu Gly Glu Asn Asn Cys Leu Leu His Pro Met Ser Gln His Gly Ile Glu Asp Pro Glu Lys Glu Val Leu Glu Trp Arg Phe Asp Ser Lys Leu Ala Phe His His Val Ala Arg Glu Leu His Pro Glu Tyr Tyr Lys Asp Cys (SEQ ID NO:12). Therefore, this exemplified Nef protein, Opt tPA-Nef, contains both a tPA leader sequence as well as deleting the myristylation site of Gly-2A DNA molecule encoding HIV-1 Nef from the HIV-1 jfrl isolate wherein the codons are optimized for expression in a mammalian system such as a human.

In another specific embodiment of the present invention, a DNA molecule is disclosed which encodes optimized HIV-1 Nef wherein the open reading frame of a recombinant adenoviral HIV vaccine encodes for modifications at the amino terminal myristylation site (Gly-2 to Ala-2) and substitution of the Leu-174-Leu-175 dileucine motif to Ala-174-Ala-175. This open reading frame is herein described as opt nef (G2A,LLAA) and is disclosed as SEQ ID NO:13, which comprises an initiating methionine residue at nucleotides 12-14 and a "TAA" stop codon from nucleotides 660-662. The nucleotide sequence of this codon optimized version of HIV-1 jrfl nef gene with the above mentioned modifications is disclosed herein as SEQ ID NO:13, as follows:

GATCTGCCAC CATGGCCGGC AAGTGGTCCA AGAGGTCCGT GCCCGGCTGG TCCACCGTGA
GGGAGAGGAT GAGGAGGGCC GAGCCCGCCG CCGACAGGGT GAGGAGGACC GAGCCCGCCG
CCGTGGGCGT GGGCGCCGTG TCCAGGGACC TGGAGAAGCA CGGCGCCATC ACCTCCTCCA
ACACCGCCGC CACCAACGCC GACTGCGCCT GGCTGGAGGC CCAGGAGGAC GAGGAGGTGG
GCTTCCCCGT GAGGCCCCAG GTGCCCCTGA GGCCCATGAC CTACAAGGGC GCCGTGGACC
TGTCCCACTT CCTGAAGGAG AAGGGCGGCC TGGAGGGCCT GATCCACTCC CAGAAGAGGC
AGGACATCCT GGACCTGTGG GTGTACCACA CCCAGGGCTA CTTCCCCGAC TGGCAGAACT
ACACCCCCGG CCCCGGCATC AGGTTCCCCC TGACCTTCGG CTGGTGCTTC AAGCTGGTGC
CCGTGGAGCC CGAGAAGGTG GAGGAGGCCA ACGAGGGCGA GAACAACTGC GCCGCCCACC
CCATGTCCCA GCACGGCATC GAGGACCCCG AGAAGGAGGT GCTGGAGTGG AGGTTCGACT
CCAAGCTGGC CTTCCACCAC GTGGCCAGGG AGCTGCACCC CGAGTACTAC AAGGACTGCT
AAAGCCCGGG C (SEQ ID NO:13).

The open reading frame of SEQ ID NO:13 encodes Nef (G2A,LLAA), disclosed herein as SEQ ID NO:14, as follows:

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Met Ala Gly Lys Trp Ser Lys Arg Ser Val Pro Gly Trp Ser Thr Val 15 Arg Glu Arg Met Arg Arg Ala Glu Pro Ala Ala Asp Arg Val Arg Arg Thr Glu Pro Ala Ala Val Gly Val Gly Ala Val Ser Arg Asp Leu Glu Lys His Gly Ala Ile Thr Ser Ser Asn Thr Ala Ala Thr Asn Ala Asp Cys Ala Trp Leu Glu Ala Gln Glu Asp Glu Glu Val Gly Phe Pro Val Arg Pro Gln Val Pro Leu Arg Pro Met Thr Tyr Lys Gly Ala Val Asp 20 Leu Ser His Phe Leu Lys Glu Lys Gly Gly Leu Glu Gly Leu Ile His Ser Gln Lys Arg Gln Asp Ile Leu Asp Leu Trp Val Tyr His Thr Gln Gly Tyr Phe Pro Asp Trp Gln Asn Tyr Thr Pro Gly Pro Gly Ile Arg Phe Pro Leu Thr Phe Gly Trp Cys Phe Lys Leu Val Pro Val Glu Pro Glu Lys Val Glu Glu Ala Asn Glu Gly Glu Asn Asn Cys Ala Ala His 25 Pro Met Ser Gln His Gly Ile Glu Asp Pro Glu Lys Glu Val Leu Glu Trp Arg Phe Asp Ser Lys Leu Ala Phe His His Val Ala Arg Glu Leu His Pro Glu Tyr Tyr Lys Asp Cys Ser (SEQ ID NO:14).

An additional embodiment of the present invention relates to another DNA molecule encoding optimized HIV-1 Nef wherein the amino terminal myristylation site and dileucine motif have been deleted, as well as comprising a tPA leader peptide. This DNA molecule, opt tpanef (LLAA) comprises an open reading frame which encodes a Nef protein containing a tPA leader sequence fused to amino acid residue 6-216 of HIV-1 Nef (jfrl), wherein Leu-174 and Leu-175 are substituted with Ala-174 and Ala-175 (Ala-195 and Ala-196 in this tPA-based fusion protein). The nucleotide

sequence comprising the open reading frame of opt tpanef (LLAA) is disclosed herein as SEQ ID NO:15, as shown below:

CATGGATGCA ATGAAGAGAG GGCTCTGCTG TGTGCTGCTG CTGTGTGGAG CAGTCTTCGT TTCGCCCAGC GAGATCTCCT CCAAGAGGTC CGTGCCCGGC TGGTCCACCG TGAGGGAGAG GATGAGGAGG GCCGAGCCCG CCGCCGACAG GGTGAGGAGG ACCGAGCCCG CCGCCGTGGG CGTGGGCGCC GTGTCCAGGG ACCTGGAGAA GCACGGCGCC ATCACCTCCT CCAACACCGC CGCCACCAAC GCCGACTGCG CCTGGCTGGA GGCCCAGGAG GACGAGGAGG TGGGCTTCCC CGTGAGGCCC CAGGTGCCCC TGAGGCCCAT GACCTACAAG GGCGCCGTGG ACCTGTCCCA CTTCCTGAAG GAGAAGGGCG GCCTGAGGG CCTGATCCAC TCCCAGAAGA GGCAGGACAT CCTGGACCTG TGGGTGTACC ACACCCAGGG CTACTTCCCC GACTGGCAGA ACTACACCCC CGGCCCCGC ATCAGGTTCC CCCTGACCTT CGGCTGGTGC TTCAAGCTGG TGCCCGTGGA GCCCGAGAAG GTGGAGGAGG CCAACGAGGG CGAGAACAAC TGCGCCGCCC ACCCCATGTC CCAGCACGC ATCGAGGACC CCGAGAAGGA GGTGCTGGAG TGGAGGTTCG ACTCCAAGCT GGCCTTCCAC CACGTGGCCA GGGAGCTGCA CCCCGAGTAC TACAAGGACT GCTAAAGCCC (SEQ ID NO:15).

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The open reading frame of SEQ ID NO:7 encoding tPA-Nef (LLAA), disclosed herein as SEQ ID NO:16, is as follows:

Met Asp Ala Met Lys Arg Gly Leu Cys Cys Val Leu Leu Cys Gly Ala Val Phe Val Ser Pro Ser Glu Ile Ser Ser Lys Arg Ser Val Pro 20 Gly Trp Ser Thr Val Arg Glu Arg Met Arg Arg Ala Glu Pro Ala Ala Asp Arg Val Arg Arg Thr Glu Pro Ala Ala Val Gly Val Gly Ala Val Ser Arg Asp Leu Glu Lys His Gly Ala Ile Thr Ser Ser Asn Thr Ala Ala Thr Asn Ala Asp Cys Ala Trp Leu Glu Ala Gln Glu Asp Glu Glu Val Gly Phe Pro Val Arg Pro Gln Val Pro Leu Arg Pro Met Thr Tyr Lys Gly Ala Val Asp Leu Ser His Phe Leu Lys Glu Lys Gly Gly Leu Glu Gly Leu Ile His Ser Gln Lys Arg Gln Asp Ile Leu Asp Leu Trp Val Tyr His Thr Gln Gly Tyr Phe Pro Asp Trp Gln Asn Tyr Thr Pro Gly Pro Gly Ile Arg Phe Pro Leu Thr Phe Gly Trp Cys Phe Lys Leu Val Pro Val Glu Pro Glu Lys Val Glu Glu Ala Asn Glu Gly Glu Asn Asn Cys Ala Ala His Pro Met Ser Gln His Gly Ile Glu Asp Pro Glu Lys Glu Val Leu Glu Trp Arg Phe Asp Ser Lys Leu Ala Phe His His Val Ala Arg Glu Leu His Pro Glu Tyr Tyr Lys Asp Cys (SEQ ID NO:16). An adenoviral vector of the present invention may comprise a DNA sequence, regardless of codon usage, which expresses a wild type or modified Nef protein as described herein, including but not limited to modified Nef proteins which comprise a deletion or substitution of Gly 2, a deletion of substitution of Leu 174 and Leu 175

and/or inclusion of a leader sequence. Therefore, partial or fully codon optimized DNA vaccine expression vector constructs are preferred since such constructs should result in increased host expression. However, it is within the scope of the present invention to utilize "non-codon optimized" versions of the constructs disclosed herein, especially modified versions of HIV Nef which are shown to promote a substantial cellular immune response subsequent to host administration.

Figure 20A-C show nucleotide sequences at junctions between nef coding sequence and plasmid backbone of nef expression vectors V1Jns/nef (Figure 20A), V1Jns/nef(G2A,LLAA) (Figure 20B), V1Jns/tpanef (Figure 20C) and V1Jns/tpanef(LLAA) (Figure 20C, also). 5' and 3' flanking sequences of codon optimized nef or codon optimized nef mutant genes are indicated by bold/italic letters; nef and nef mutant coding sequences are indicated by plain letters. Also indicated (as underlined) are the restriction endonuclease sites involved in construction of respective nef expression vectors. V1Jns/tpanef and V1Jns/tpanef(LLAA) have identical sequences at the junctions.

Figure 21 shows a schematic presentation of nef and nef derivatives. Amino acid residues involved in Nef derivatives are presented. Glycine 2 and Leucine174 and 175 are the sites involved in myristylation and dileucine motif, respectively.

20 EXAMPLE 19

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MRKAd5Pol Construction and Virus Rescue

Steps performed in the construction of the vectors, including the pre-adenovirus plasmid - Key steps performed in the construction of the vectors, including the pre-adenovirus plasmid denoted MRKAd5pol, is depicted in Figure 22. Briefly, the adenoviral shuttle vector for the full-length inactivated HIV-1 pol gene is as follows. The vector MRKpdelE1(Pac/pIX/pack450)+CMVmin+BGHpA(str.) is a derivative of the shuttle vector used in the construction of the MRKAd5gag adenoviral pre-plasmid. The vector contains an expression cassette with the hCMV promoter (no intronA) and the bovine growth hormone polyadenylation signal. The expression unit has been inserted into the shuttle vector such that insertion of the gene of choice at a unique BgIII site will ensure the direction of transcription of the transgene will be Ad5 E1 parallel when inserted into the MRKpAd5(E1-/E3+)Cla1 (or MRKpAdHVE3) preplasmid. The vector, similar to the original shuttle vector contains the Pac1 site, extension to the packaging signal region, and extension to the pIX gene. The synthetic full-length codon-optimized HIV-1 pol gene was isolated directly from the plasmid pV1Ins-HIV-pol-inact(opt). Digestion of this plasmid with BgI II releases the pol

gene intact (comprising a codon optimized IA pol sequence as disclosed in SEQ ID NO:3). The pol fragment was gel purified and ligated into the MRKpdelE1(Pac/pIX/pack450)+CMVmin+BGHpA(str.) shuttle vector at the BgIII site. The clones were checked for the correct orientation of the gene by using restriction enzymes DraIII/Not1. A positive clone was isolated and named MRKpdel+hCMVmin+FL-pol+bGHpA(s). The genetic structure of this plasmid was verified by PCR, restriction enzyme and DNA sequencing. The pre-adenovirus plasmid was constructed as follows. Shuttle plasmid MRKpdel+hCMVmin+FLpol+bGHpA(S) was digested with restriction enzymes Pac1 and Bst1107 I (or its isoschizomer, BstZ107 I) and then co-transformed into E. coli strain BJ5183 with linearized (Cla1 digested) adenoviral backbone plasmid, MRKpAd(E1-/E3+)Cla1. The resulting pre-plasmid originally named MRKpAd+hCMVmin+FLpol+bGHpA(S)E3+ is now referred to as "pMRKAd5pol". The genetic structure of the resulting pMRKAd5pol was verified by PCR, restriction enzyme and DNA sequence analysis. The vectors were transformed into competent E. coli XL-1 Blue for preparative production. The recovered plasmid was verified by restriction enzyme digestion and DNA sequence analysis, and by expression of the pol transgene in transient transfection cell culture. The complete nucleotide sequence of this pMRKAd5HIV-1pol adenoviral vector is shown in Figure 26 A-AO.

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Generation of research-grade recombinant adenovirus - The pre-adenovirus plasmid, pMRKAd5pol, was rescued as infectious virions in PER.C6® adherent monolayer cell culture. To rescue infectious virus, 12 μ g of pMRKAd5pol was digested with restriction enzyme PacI (New England Biolabs) and 3.3 μ g was transfected per 6 cm dish of PER.C6® cells using the calcium phosphate coprecipitation technique (Cell Phect Transfection Kit, Amersham Pharmacia Biotech Inc.). PacI digestion releases the viral genome from plasmid sequences allowing viral replication to occur after entry into PER.C6® cells. Infected cells and media were harvested 6 -10 days post-transfection, after complete viral cytopathic effect (CPE) was observed. Infected cells and media were stored at \leq -60°C. This pol containing recombinant adenovirus is referred to herein as "MRKAd5pol". This recombinant adenovirus expresses an inactivated HIV-1 Pol protein as shown in SEQ ID NO:6.

EXAMPLE 20

MRKAd5Nef Construction and Virus Rescue

Construction of vector: shuttle plasmid and pre-adenovirus plasmid - Key steps performed in the construction of the vectors, including the pre-adenovirus

plasmid denoted MRKAd5nef, is depicted in Figure 23. Briefly, as shown in Example 19 above, the vector

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MRKpdelE1(Pac/pIX/pack450)+CMVmin+BGHpA(str.) is the shuttle vector used in the construction of the MRKAd5gag adenoviral pre-plasmid. It has been modified to contain the *Pac*1 site, extension to the packaging signal region, and extension to the pIX gene. It contains an expression cassette with the hCMV promoter (no intronA) and the bovine growth hormone polyadenylation signal. The expression unit has been inserted into the shuttle vector such that insertion of the gene of choice at a unique *Bgl*11 site will ensure the direction of transcription of the transgene will be Ad5 E1 parallel when inserted into the MRKpAd5(E1-/E3+)Cla1 pre-plasmid. The synthetic full-length codon-optimized HIV-1 nef gene was isolated directly from the plasmid pV1Jns/nef (G2A,LLAA). Digestion of this plasmid with *Bgl*11 releases the pol gene intact, which comprises the nucleotide sequence as disclosed in SEQ ID NO:13. The nef fragment was gel purified and ligated into the

MRKpdelE1+CMVmin+BGHpA(str.) shuttle vector at the Bgl11 site. The clones were checked for correction orientation of the gene by using restriction enzyme Scal. A positive clone was isolated and named MRKpdelE1hCMVminFL-nefBGHpA(s). The genetic structure of this plasmid was verified by PCR, restriction enzyme and DNA sequencing. The pre-adenovirus plasmid was constructed as follows. Shuttle plasmid MRKpdelE1hCMVminFL-nefBGHpA(s) was digested with restriction enzymes Pac1 and Bst1107 I (or its isoschizomer, BstZ107 I) and then co-transformed into E. coli strain BJ5183 with linearized (Cla1 digested) adenoviral backbone plasmid, MRKpAd(E1/E3+)Cla1. The resulting pre-plasmid originally named MRKpdelE1hCMVminFL-nefBGHpA(s) is now referred to as "pMRKAd5nef". The genetic structure of the resulting pMRKAd5nef was verified by PCR, restriction enzyme and DNA sequence analysis. The vectors were transformed into competent E. coli XL-1 Blue for preparative production. The recovered plasmid was verified by restriction enzyme digestion and DNA sequence analysis, and by expression of the nef transgene in transient transfection cell culture. The complete nucleotide sequence of this pMRKAd5HIV-1nef adenoviral vector is shown in Figure 27A-AM.

Generation of research-grade recombinant adenovirus - The pre-adenovirus plasmid, pMRKAd5nef, was rescued as infectious virions in PER.C6® adherent monolayer cell culture. To rescue infectious virus, 12 μg of pMRKAdnef was digested with restriction enzyme Pac1 (New England Biolabs) and 3.3 μg was transfected per 6 cm dish of PER.C6® cells using the calcium phosphate co-precipitation technique (Cell Phect Transfection Kit, Amersham Pharmacia Biotech

Inc.). Pac1 digestion releases the viral genome from plasmid sequences allowing viral replication to occur after entry into PER.C6[®]cells. Infected cells and media were harvested 6-10 days post-transfection, after complete viral cytopathic effect (CPE) was observed. Infected cells and media were stored at \leq -60°C. This nef containing recombinant adenovirus is now referred to as "MRKAd5nef".

EXAMPLE 21

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Construction of Murine CMV Promoter Containing Shuttle Vectors for Inactivated Pol and Nef/G2A,LLAA

The murine CMV (mCMV) was amplified from the plasmid pMH4 (supplied by Frank Graham, McMaster University) using the primer set: mCMV (Not I) Forward: 5'-ATA AGA ATG CGG CCG CCA TAT ACT GAG TCA TTA GG-3' (SEQ ID NO: 20); mCMV (Bgl II)Reverse: 5'-AAG GAA GAT CTA CCG ACG CTG GTC GCG CCT C-3' (SEQ ID NO:21). The underlined nucleotides represent the Not I and the Bgl II sites respectively for each primer. This PCR amplicon was used for the construction of the mCMV shuttle vector containing the transgene in the E1 parallel orientation. The hCMV promoter was removed from the original shuttle vector (containing the hCMV-gag-bGHpA transgene in the E1 parallel orientation) by digestion with Not I and Bgl II. The mCMV promoter (Not I/Bgl II digested PCR product) was inserted into the shuttle vector in a directional manner. The shuttle vector was then digested with $Bgl \ \Pi$ and the gag reporter gene ($Bgl \ \Pi$ fragment) was re-inserted back into the shuttle vector. Several clones were screened for correct orientation of the reporter gene. For the construction of the mCMV-gag in the E1 antiparallel orientation, the mCMV promoter was amplified from the plasmid pMH4 using the following primer set: mCMV (Asc I) Forward: 5'- ATA AGA ATG GCG \underline{CGC} CAT ATA CTG AGT CAT TAG G (SEQ ID NO:22); mCMV (Bgl II) Reverse: 5' AAG GAA GAT CTA CCG ACG CTG GTC GCG CCT C (SEQ ID NO:23). The underlined nucleotides represent the Asc I and Bgl II sites, respectively for each primer. The shuttle vector containing the hCMV-gag transgene in the E1 antiparallel orientation was digested with Asc1 and Bgl11 to remove the hCMV-gag portion of the transgene. The mCMV promoter (Asc1/Bgl11 digested PCR product) was inserted into the shuttle vector in a directional manner. The vector was then digested with Bgl11 and the gag reporter gene (Bgl11 fragment) was re-inserted. Several clones were screened for correct orientation of the reporter gene. For each of the full length IA pol and full length nef/G2A,LLAA genes, cloning was performed using the unique

 $Bgl \ \Pi$ site within the mCMV-bGHpA shuttle vector. The pol and nef genes were excised from their respective pV1Jns plasmids by $Bgl \ \Pi$ digestion.

EXAMPLE 22

Construction of mCMV Full Length Inactivated Pol and Full Length nef/G2A.LLAA Adenovectors

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Each of these transgenes of Example 21 were inserted into the modified shuttle vector in both the E1 parallel and E1 anti-parallel orientations. *Pac1* and *BstZ110I* digestion of each shuttle vector was performed and each specific transgene fragment containing the flanking Ad5 sequences was isolated and co-transformed with *Cla I* digested MRKpAd5(E3+) or MRKpAd5(E3-) adenovector plasmids via bacterial homologous recombination in BJ5183 *E. coli* cells. Recombinant preplasmid adenovectors containing the various transgenes in both the E3- and E3+ versions (and in the E1 parallel and E1 antiparallel orientations) were subsequently prepared in large scale following transformation into XL-1 Blue *E. coli* cells and analyzed by restriction analysis and sequencing.

EXAMPLE 23

Construction of hCMV-tpa-nef (LLAA) Adenovector

The tpa-nef gene was amplified out from GMP grade pV1Jns-tpanef (LLAA) vector
using the primer sets: Tpanef (BamHI) F 5'-ATT GGA TCC ATG GAT GCA ATG

AAG AGA GGG (SEQ ID 24); Tpanef (BamHI) R 5'-ATA GGA TCC TTA GCA
GTC CTT GTA GTA CTC G (SEQ ID NO:25). The resulting PCR product was
digested with BamHI, gel purified and cloned into the Bgl II site of MRKAd5CMVbGHpA shuttle vector (Bgl II digested and calf intestinal phosphatase treated).
Clones containing the tpanef (LLAA) gene (see SEQ ID NO:15 for complet coding
region) in the correct orientation with respect to the hCMV promoter were selected
following Sca I digestion. The resulting MRKAd5tpanef shuttle vector was digested
with Pac I and Bst Z1101 and cloned into the E3+ MRKAd5 adenovector via bacterial
homologous recombination techniques.

EXAMPLE 24

Immunogenicity of MRKAd5pol and MRKAd5nef Vaccine

Materials and Methods - Rodent Immunization - Groups of N=10 BALB/c

mice were immunized i.m. with the following vectors: (1) MRKAd5hCMV-IApol

(E3+) at either 10^7 vp and 10^9 vp; and (2) MRKAd5hCMV-IApol (E3-) at either

10^7 vp and 10^9 vp. At 7 weeks post dose, 5 of the 10 mice per cohort were boosted with the same vector and dose they initially received. At 3 weeks post the second does, sera and spleens were collected from all the animals for RT ELISA and IFNg ELIspot analyses, respectively. For all rodent immunizations, the Ad5 vectors were diluted in 5 mM Tris, 5% sucrose, 75 mM NaCl, 1 mM MgCl2, 0.005% polysorbate 80, pH 8.0. The total dose was injected to both quadricep muscles in 50 µL aliquots using a 0.3-mL insulin syringe with 28-1/2G needles (Becton-Dickinson, Franklin Lakes, NJ).

Groups of N=10 C57/BL6 mice were immunized i.m. with the following vectors: (1) MRKAd5hCMV-nef(G2A,LLAA) (E3+) at either 10^7 vp and 10^9 vp; (2) MRKAd5mCMV-nef(G2A,LLAA) (E3+) at either 10^7 vp and 10^9 vp; and (3) MRKAd5mCMV-tpanef(LLAA) (E3+) at either 10^7 vp and 10^9 vp. At 7 weeks post dose, 5 of the 10 mice per cohort were boosted with the same vector and dose they initially received. At 3 weeks post the second does, sera and spleens were collected from all the animals for RT ELISA and IFNg ELIspot analyses, respectively.

Non-human Primate immunization - Cohorts of 3 rhesus macaques (2-3 kg) were vaccinated with the following Ad vectors: (1) MRKAd5hCMV-IApol (E3+) at either 10^9 vp and 10^11 vp dose; and (2) MRKAd5hCMV-IApol (E3-) at either 10^9 vp and 10^11 vp; (3) MRKAd5hCMV-nef(G2A,LLAA) (E3+) at either 10^9 vp and 10^11 vp; and (4) MRKAd5mCMV-nef(G2A,LLAA) (E3+) at either 10^9 vp and 10^11 vp. The vaccine was administered to chemically restrained monkeys (10 mg/kg ketamine) by needle injection of two 0.5 mL aliquots of the Ad vectors (in 5 mM Tris, 5% sucrose, 75 mM NaCl, 1 mM MgCl₂, 0.005% polysorbate 80, pH 8.0) into both deltoid muscles. The animals were immunized twice at a 4 week interval (T=0, 4 weeks).

Murine anti-RT and anti-nef ELISA - Anti-RT titers were obtained following standard secondary antibody-based ELISA. Maxisorp plates (NUNC, Rochester; NY) were coated by overnight incubation with 100 μL of 1 μg/mL HIV-1 RT protein (Advanced Biotechnologies, Columbia, MD) in PBS. For anti-nef ELISA, 100 uL of 1 ug/mL HIV-1 nef (Advanced Biotechnologies, Columbia, MD) was used to coat the plates. The plates were washed with PBS/0.05% Tween 20 using Titertek MAP instrument (Hunstville, AL) and incubated for 2 h with 200 μL/well of blocking solution (PBS/0.05% tween/1% BSA). An initial serum dilution of 100-fold was performed followed by 4-fold serial dilution. 100-μL aliquots of serially diluted samples were added per well and incubated for 2 h at room temperature. The plates

were washed and 100 μL of 1/1000-diluted HRP-rabbit anti-mouse IgG (ZYMED, San Francisco, CA) were added with 1 h incubation. The plates were washed thoroughly and soaked with 100 μL 1,2-phenylenediamine dihydrochloride/hydrogen peroxide (DAKO, Norway) solution for 15 min. The reaction was quenched by adding 100 μL of 0.5M H₂SO4 per well. OD₄₉₂ readings were recorded using Titertek Multiskan MCC/340 with S20 stacker. Endpoint titers were defined as the highest serum dilution that resulted in an absorbance value of greater than or equal to 0.1 OD₄₉₂ (2.5 times the background value).

Non-human primate and murine ELIspot assays - The enzyme-linked immuno-spot (ELISpot) assay was utilized to enumerate antigen-specific INFy-secreting cells from mouse spleens (Miyahira, et al.1995, J. Immunol. Methods 181:45-54) or macaque PBMCs. Mouse spleens were pooled from 5 mice/cohort and single cell suspensions were prepared at 5×10^6 /mL in complete RPMI media (RPMI1640, 10% FBS, 2mM L-glutamine, 100U/mL Penicillin, 100 u/mL streptomycin, 10 mM Hepes, 50 uM β -ME). Rhesus PBMCs were prepared from 8-15 mL of heparinized blood following standard Ficoll gradient separation (Coligan, et al, 1998, Current Protocols in Immunology. John Wiley & Sons, Inc.). Multiscreen opaque plates (Millipore, France) were coated with 100 μ L/well of either 5 μ g/mL purified rat anti-mouse IFN- γ IgG1, clone R4-6A2 (Pharmingen, San Diego, CA), or 15 ug/mL mouse anti-human IFN- γ IgG2a (Cat. No. 1598-00, R&D Systems, Minneapolis, MN) in PBS at 4°C overnight for murine or monkey assays, respectively. The plates were washed with PBS/penicillin/streptomycin and blocked with 200 μ L/well of complete RPMI media for 37 °C for at least 2 h.

To each well, 50 μL of cell samples (4-5x10⁵ cells per well) and 50 μL of the antigen solution were added. To the control well, 50 μL of the media containing DMSO were added; for specific responses, either selected peptides or peptide pools (4 ug/mL per peptide final concentration) were added. For BALB/c mice immunized with the pol constructs, stimulation was conducted using a pool of CD4⁺-epitope containing 20-mer peptides (aa21-40, aa411-430, aa641-660, aa731-750, aa771-790) or a pool of CD8⁺-epitope containing peptides (aa201-220, aa311-330, aa781-800). For C57/BL6 mice immunized with the nef construct, either aa51-70 (CD8⁺ T cell epitope) or aa81-100 (CD4⁺) peptide derived from the nef sequence was added for specific stimulation. In monkeys, the responses against pol were evaluated using two pools (L and R) of 20-aa peptides that encompass the entire pol sequence and overlap by 10 amino acids. In monkeys vaccinated with the nef constructs, a single pool containing 20-mer peptides covering the entire HIV-1 nef sequence and overlapping

by 10 aa was used. Each sample/antigen mixture was performed in triplicate wells for murine samples or in duplicate wells for rhesus PBMCs. Plates were incubated at 37° C, 5% CO₂, 90% humidity for 20-24 h. The plates were washed with PBS/0.05% Tween 20 and incubated with 100 μ L/well of either 1.25 μ g/mL biotin-conjugated rat anti-mouse IFN- γ mAb, clone XMG1.2 (Pharmingen) or of 0.1 μ g/mL biotinylated anti-human IFN-gamma goat polyclonal antibody (R&D Systems) at 4°C overnight. The plates were washed and incubated with 100 μ L/well 1/2500 dilution of strepavidin-alkaline phosphatase conjugate (Pharmingen) in PBS/0.005% Tween/5% FBS for 30 min at 37 °C. Spots were developed by incubating with 100 μ L/well 1-step NBT/BCIP (Pierce Chemicals) for 6-10 min. The plates were washed with water and allowed to air dry. The number of spots in each well was determined using a dissecting microscope and the data normalized to 10^{6} cell input.

Non-human Primate anti-RT ELISA - The pol-specific antibodies in the monkeys were measured in a competitive RT EIA assay, wherein sample activity is determined by the ability to block RT antigen from binding to coating antibody on the plate well. Briefly, Maxisorp plates were coated with saturating amounts of pol positive human serum (#97111234). 250 uL of each sample is incubated with 15 uL of 266 ng/mL RT recombinant protein (in RCM 563, 1% BSA, 0.1% tween, 0.1% NaN₃) and 20 uL of lysis buffer (Coulter p24 antigen assay kit) for 15 min at room temperature. Similar mixtures are prepared using serially diluted samples of a standard and a negative control which defines maximum RT binding. 200 uL/well of each sample and standard were added to the washed plate and the plate incubated 16-24 h at room temperature. Bound RT is quantified following the procedures described in Coulter p24 assay kit and reported in milliMerck units per mL arbitrarily defined by the chosen standard.

Results - Rodent Studies - BALB/c mice (n=5 mice/cohort) were immunized once or twice with varying doses of MRKAd5hCMV-IApol(E3+) and MRKAd5hCMV-IApol(E3-). At 3 weeks after the second dose, Anti-pol IgG levels were determined by an ELISA assay using RT as a surrogate antigen. Cellular response were quantified via IFNy ELISpot assay against pools of pol-epitope containing peptides. The results of these assays are summarized in Table 10. The results indicate that the mouse vaccinees exhibited detectable anti-RT IgGs with an adenovector dose as low as 10^7 vp. The humoral responses are highly dose-dependent and are boostable with a second immunization. One or two doses of either pol vectors elicit high frequencies of antigen-specific CD4+ and CD8+T cells; the responses are weakly dose-dependent but are boostable with a second immunization.

Table 10. Immunogenicity of MRKAd5pol Vectors in BALB/c mice.

	TO: Minney-Control	, 02									
				Ап	ti-RT IgG Tite	rs"	S	FC/10^6 cell	s°		
Group	Vaccine	Dose	No. of Doses	GMT	+SE	-SE	Medium	CD4+ peptide pool	CD8+ peptide pool		
1	MRKAd5hCMVFLpol (E3+)	10^7 vp	2 1	310419 919	301785 372	153020 265	1(1) 1(1)	75(4) 72(9)	2313(67) 533(41)		
2	MRKAd5hCMVFLpol (E3+)	10^9 vp	2 1	1838400 ^b 713155	0 528520	0 303555	2(2) 1(1)	114(9) 48(7)	2063(182) 733(89)		
3	MRKAd5hCMVFLpol (E3-)	10^7 vp	2	310419 6400	386218 14013	172097 4393	0(0) 10(8)	223(7) 141(21)	2607(27) 409(28)		
4	MRKAd5hCMVFLpol (E3-)	10^9 vp	2	1838400 ^b 1241675 ^b	0 396725	0 300681	1(1) 0(0)	160(13) 39(13)	2385(11) 833(83)		
5	Naïve	none	none	57	9	7	9(2)	11(4)	10(1)		

^{*}GMT, geometric mean titer of the cohort of 5 mice; SE, standard error of the gemetric mean

5 C57/BL6 mice were immunized once or twice with varying doses of MRKAd5hCMV-nef(G2A,LLAA) (E3+), MRKAd5mCMV-nef(G2A,LLAA) (E3+) at either 10^7 vp and(3) MRKAd5mCMV-tpanef(LLAA) (E3+) at either 10^7 vp and 10^9 vp. The immune response were analyzed using similar protocols and the results are listed in Table 11. While anti-nef IgG responses could not be detected in this model system with any of the constructs, there are strong indications of a cellular immunity generated against nef using the ELIspot assay.

Table 11. Immunogenicity of MRKAd5nef Vectors in C57/BL6 mice.

				Ar	iti-nef lgG Tite	ers"	s	FC/10^6 cell	5 ^b
Group	Vaccine	Dose	No. of Doses	GMT	+SE	-SE	Medium	aa51-70 CD8+	aa81-100 CD4+
1	MRKAd5hCMVFLnef (E3+)	10^7 vp	2	174 132	70 42	50 32	1(1) 0(0)	23(1) 0(0)	1(1) 0(0)
2	MRKAd5hCMVFLnef (E3+)	10 ^9 vp	2	174 132	70 42	50 32	0(0) 1(1)	61(7) 62(7)	4(2) 3(1)
3	MRKAd5mCMVFLnef (E3+)	10^7 vp	2 1 ·	132 115	42 46	32 33	3(1) 3(2)	15(5) 3(2)	5(2) 4(2)
4	MRKAd5mCMVFLnef (E3+)	10^9 vp	2	132 132	42 42	32 32	4(2) 2(1)	83(13) 29(2)	5(1) 4(0)
5	MRKAd5mCMVtpanef(E3+)	10^7 vp	2 1	132 100	42 0	32 0	3(2) 3(1)	14(2) 13(4)	5(1) 10(3)
6	MRKAd5mCMVlpanef(E3+)	10^9 vp	2	230 115	170 46	98 33	3(2) 7(1)	145(29) 151(14)	4(0) 10(0)
7	Naïve	поле	none	152	78	52 ·	· 21(2)	- 18(6)	26(3)

*GMT, geometric mean titler of the cohort of 5 mice; SE, standard error of the gemetric mean

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Monkey Studies - Cohorts of 3 rhesus macaques were immunized with 2 doses of MRKAd5hCMV-IApol(E3+) and MRKAd5hCMV-IApol(E3-). The number of antigen-specific T cells (per million PBMCs) were enumerated using one of two

Near or at the upper limit of the serial dilution; hence, could be greater than this value

[&]quot;No. of Spot-forming Cells per million splechoytes; mean values of triplicates are reported along with standard errors in parenthesis.

No. of spot-forming cells per million spiechoytes; mean values of triplicates are reported along with standard errors in parenthesis.

peptide pools (L and R) that cover the entire pol sequence; the results are listed in Table 12. Moderate-to-strong T cell responses were detected in the vaccinees using either constructs even at a low dose of 10^9 vp. Longitudinal analyses of the anti-RT antibody titers in the animals suggest that the pol transgene product is expressed efficiently to elicit a humoral response (Table 13). It would appear that generally higher immune responses were observed in animals that received the E3- construct compared to the E3+ virus.

Table 12. Pol-specific T Cell Responses in MRKAd5pol Immunized Rhesus

10	Macaques.
10	TYTACACHUCS.

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Vaccine (T=0,4 wks)	Monk #		Prebleed			T=4			T=7			T=16	
		Mock	Pol L	Pol R	Mock	Pol L	Pol R	Mock	Pol L	Pol R	Mock	Pol L	Pol R
MRKAd5hCMV-IApod(E3+)	99C100	1	0	0	1	38	31	0	52	146	0	49	715
10^11 VD	99C215	1	2	2	10	98	249	1 1	109	305	22	88	250
10:11 VP	99D201	5	5	4	6	149	85	0	40	35	0	35	18
MRKAd5hCMV-IApol(E3+)	99D212	0	2	0	4	331	114	0	58	14	0	6	6
10/9 VD	99D180	0	4	2	0	19	192	4	38	156	5	38	108
	99C201	8	5	21	6	62	62	0	18	32	۱ ۱	14	65
MRKAd5hCMV-IApd(E3-)	99D239	5	2	2	20	82	172	1	66	114	9	21	40
10411 VD	99C186	4	12	6	5	120	421	2	271	489	16	875	530
	99C084	1	8	9	8	84	464	٥	14	238	1	24	264
MRKAd5hCMV-IApod(E3-)	CC7C	10	10	8	12	724	745	4	322	376	4	188	176
10'9 VD	CDIG	2	0	1	5	474	468	0	232	212	0	101	121
	0011	6	6	12	10	98	110	5	60	80	8	25	34
Nave	083Q	nd	nd	nd	nd	nd	nd	4	2	2	2	1	2

nd, not determined Reported are SFC per million PBMCs; mean of duplicate wells.

Table 13. Anti-RT Ig Levels in MRKAd5pol Immunized macaques.

RT ANTIBODY ASSAY TITERS IN MMU/	mL			
Vaccine/Monkey T ag	1=4	T =7	T=12	T=16
MRKAd5hCMV-IApol(E3+), 10^11 vp				
99C100	61	1999	5928	4768
99C215	81	1541	2356	2767
99D201	53	336	539	387
MRKAd5hCMV-IApol(E3+), 10^9 vp				
99D212	10	40	49	68
99D180	<10	36	79	93
99C201	<10	37	71	76
MRKAd5hCMV-IApol(E3-), 10^11 vp				
99D239	44	460	1234	1015
99C186	21	· 233 ·	480	345
990084	235	2637	2858_	1626
MRKAd5hCMV-IApol(E3-), 10^9 vp				
CC7C	32	175	306	235_
Φ16	20	140	273	419
Q11	15	112	149	237

When rhesus macaques were immunized i.m. with two doses of MRKAd5nef constructs, vigorous T cell responses ranging from 100 to as high as 1100 per million were observed in 8 of 12 vaccinees (Table 14). The efficacies of the mCMV- and hCMV- driven nef constructs are comparable on the basis of the data generated thus far.

Table 14. Nef-specific T cell Responses in MRKAd5nef Immunized Rhesus

Macaques.

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Vaccine (T=0,4 wks)	Monk #	Pi	е	Te	:4	T=	:7	To	16
	····	Mock	Nef	Mock	Nef	Mock	Nef	Mock	Nef
MRKAd5hCMV-nef(G2A,LLAA) (E3+)	CD2D	0	4	31	440	4	368	1	251
10^11 vp	CC7B	0	0	2	521	0	178	1	152
	CC61	2	9	31	112	0	108	11	100
MRKAd5hCMV-nef(G2A,LLAA) (E3+)	CC2K	9	9	6	52	0	35	0	15
10^9 vp	CD15	5	4	30	998	2	586	0	434
•	CD16	6	1	6	1146	0	369	1	21
MRKAd5mCMV-nef(G2A,LLAA) (E3+)	99D191	1	5	4	614	0	298	2	419
10^11 vp	99D144	4	6	5	434	0	1100	2	93
·	99C193	1	2	1	58	1	22	٥	64
MRKAd5mCMV-nef(G2A,LLAA) (E3+)	99D224	1	11	14	231	1	125	0	70
10^9 vp	99D250	В	8	4	108	0	54	0	5
•	99C120	1	6	20	299	0	92	0	79
Naîve	083Q	nd	nd	18	22	4	5	2	1

EXAMPLE 25

Comparison of Clade B vs. Clade C T Cell Responses in HIV-Infected Subjects

PBMC samples collected from two dozens of patients infected with HIV-1 in

US were tested in ELISPOT assays with peptide pools of 20-mer peptides overlapping
by 10 amino acids. Four different peptide pools were tested for cross-clade
recognition, and they were either derived from a clade B-based isolate (gag H-b; nefb) or a clade C-based isolate (gag H-c, nef-c). Data in Table 15 shows that T cells
from these patients presumably infected with clade B HIV-1 could recognize clade C
gag and nef antigens in ELISPOT assay. Correlation analysis further demonstrated
that these T cell responses against clade C gag peptide pool were about 60% of the
clade B counterpart (Figure 24), while the T cell responses against clade C nef were
about 85% of the clade B counterpart (Figure 25). These results suggest that cellular
immune responses generated in patients infected with clade B HIV-1 can recognize
gag and nef antigens derived from clade C HIV-1. These data show that a HIV
vaccine, such as a DNA or MRKAd5-based adenoviral vaccine expressing a clade B

gag and/or nef antigen will potentially have the ability to provide a prophylactic and/or therapetic advantage on a global scale.

Table 15
Responses Shown as the Number of gIFN-Secreting T Cells per Million PBMCs

subject	bleed date	gag epitope #	mock	gag H-b	gagH-c	nef-b	nef-c
		from mapping)					
#100	19-Jul-99	12	10	3950	1385	1295	1300
#101	25-Jul-99	3	15	3885	1280	na	1020
#102	25-Jul-99	4	15	1740	850	1255	1785
#104	7-Jun-99	2	5	1355	1185	na	1060
#107	11-Oct-99	2	25	3305	2795	670	870
#405	11-Jul-99	2	15	4575	3180	1700	1500
#501	19-Jul-99	2	15	1100	570	3365	3460
#505	18-Jul-99	5	10	2145	1725	1235	na
#506	28-Feb-99	2	25	150	45	400	610
#701	28-Mar-99	5	30	7620	4775	3320	2780
#709	17-May-99	3	15	2785	1945	1090	1630
#710	24-May-99	4	5	1055	1080	2210	2140

10 EXAMPLE 26

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Characterization and Production of MRKAd5pol and MRKAd5nef
Vectors in Roller Bottles

Expansion of nef and pol Adenovectors - Nef and pol CsCl purified MRKAd5 seeds were used to infect roller bottles to produce P4 virus to be used as a seed for further experiments. P4 MRKAd5 pol and nef vectors were used to infect roller bottles at an MOI 280 vp/cell, except for hCMV-tpa-nef [E3+] which was infected at an MOI of 125 due to low titers of seed obtained at P4.

Table 16 Viral particle concentrations for P5 nef and pol adenovectors

Adenovector	AEX Titer (10 ¹⁰ vp/ml culture)	AEX Titer (10 ⁴ vp/cell)	Amplification Ratio
hCMV-FL-nef [E3+]	1.1	0.9	30
mCMV-FL-nef [E3+]	2.2	2.1	75
hCMV-tpa-nef [E3+]	0.07	0.1	5
mCMV-tpa-nef [E3+]	1.3	0.9	35
hCMV-FL-pol [E3+]	2.7	2.1	75
hCMV-FL-pol [E3-]	1.9	1.3_	45

through passage seven. Cell-associated (freeze/thaw lysis) and whole broth (tritonlysis) titers obtained in all passages were very consistent. In general, MRKAd5pol is ca. 70% as productive as MRKAd5gag while MRKAd5nef is ca. 25% as productive as MRKAd5gag. Samples of P7 virus for both constructs were analyzed by V&CB by restriction digest analysis and did not show any rearrangements.

Table 17. Passage Six Viral Productivity for MRKAd5pol and MRKAd5nef

14010		Xviable (10° cells/ml), Viability (%) Infection Harvest		Cell Passage Number	AEX Titer (Cell Associated) 10 ¹⁰ vp/ml culture	Titer	Amplification Ratio	Triton Lysis Titer 10 ¹⁰ vp/ml culture
hCMV-FL-nef [B3+]	pool	1.22, 85%		62	0.8	0.7	25	1.6
	1		0.99, 62%					
	2		1.10, 72%	1	}			
hCMV-FL-pol [E3+]	pool	1.42, 89%		62	4.5	3.2	115	7.0
	. 1		1.22, 70%					
	2		1.42, 74%					

15 Table 18. Passage Seven Viral Productivity for MRKAd5pol and MRKAd5nef

15 Table		Xviable (10 Viabil Infection		Cell Passage Number	AEX Titer (Cell Associated) 10 ³⁰ vp/ml culture	Titer 10 ⁴ vp/cell	Amplification Ratio	Triton Lysis Titer 10 ¹⁰ vp/ml culture
hCMV-FL-nef [E3+]	Pool	1.33, 90%		66	1.0	0.8	29	2.1
	1		0.96, 70%					
	2		1.18, 73%	.\				
bCMV-FL-pol [E3+]	Pool	0.90*, 90%		56	4.2	4.7	168	6.5
	1		1.18, 88%					
	2		1.04, 80%	Ì				

MRKAd5nef and MRKAd5pol Viral Production Kinetics - A timecourse experiment was carried out in roller bottles to determine if the viral production kinetics of the MRKAd5pol and MRKAd5nef vectors were similar to those of
 MRKAd5gag. PER.C6® cells in roller bottle cultures were infected at an MOI of 280 vp/cells with P5 MRKAd5pol, P5 MRKAd5nef and P7 MRKAd5gag; for each adenovector, two infected bottles were sampled at 24, 36, 48, and 60 hours post infection. In addition, two bottles were left unsampled until 48 hpi when they were harvested under the Phase I process conditions. The anion-exchange HPLC viral
 particle concentrations of the freeze-thaw recovered cell associated virus at the 24, 36,

48, and 60 hpi timepoints are shown in Figure 29A-B. The QPA titers show a similar trend (data not shown).

Comparison of hCMV- and mCMV-FL-nef - As the titers obtained with the MRKAd5nef construct (hCMV-FL-nef) were lower than those obtained with MRKAd5gag or MRKAd5pol, a viral productivity comparison experiment was performed with mCMV-FL-nef. For each of the two adenovectors (hCMV- and mCMV-FL-nef), two roller bottles were infected at an MOI of 280 vp/cell with passage five clarified lysate. The macroscopic and microscopic observations of the four roller bottles were identical at the time of harvest. Analysis of the clarified lysate produced indicated a higher viral particle concentration in the bottles infected with mCMV-FL-nef, as shown in Table 19. It is stipulated that the higher productivity with mCMV promoter driven nef vector is due to lower nef expression levels in PER.C6® cells- experiments are underway at V&CB to measure nef expression levels.

Table 19. Passage Six Viral Productivity Comparison of hCMV- and mCMV-FL-nef

	[Xv (106 cells/m	l), Viability (%)	Cell Passage	AEX Titer	Titer	Amplification	Triton Lysis Titer
		Infection	Harvest	Number	10 ¹⁰ vp/ml culture	10 ⁴ vp/cell	Ratio	10 ^{to} vp/ml culture
hCMV-FL-nef	Pool	1.11,91%		60	1.5	1.4	50	2.8
(MRKAd5nef)	1		1.23,75%					
	2		1.34,74%					
mCMV-FL-nef	Pool	1.11, 91%		60	2.3	2.1	75	4.6
	1		1.49, 84%					
	2		1.18, 77%				·	

EXAMPLE 27

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Characterization and Large Scale Production of MRKAd5nef Virus in Bioreactors

Materials and Methods - The experiment of the present example was run twice under the following conditions: 36.5°C, DO 30%, pH 7.30, 150rpm agitation rate, no sparging, Life Technologies (Gibco, Invitrogen) 293 SFM II (with 6mM L-glutamine), 0.5M NaOH as base for pH control. During the first run (B20010115), two 10L stirred vessel bioreactors were inoculated with PER.C6® cells at a concentration of 0.2x106 cells/ml. Cells were grown until they reached a cell concentration of approximately 1x106 cells/ml. The cells were infected with uncloned MRKAd5nef (G2A,LLAA) at a MOI of 280 virus particles (vp)/cell. For the second batch (B20010202), the same procedure as the first run was used, except the cells

were infected with cloned MRAd5nef. During both runs, the bioreactors were harvested 48 hours post-infection. Samples were taken and virus concentrations were determined from whole broth (with triton lysis), supernatant, and cell pellets (3 X freeze/thaw) with the AEX and QPA assays. Metabolites were measured with BioProfile 250 throughout the process.

Table 20: Experimental Conditions

Temperature	36.5 °C
DO	30%
PH	7.30
Agitation	150 rpm
Sparging	None

Table 21: Virus source used for experiments.

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Run	Batch ID	Cloned/Uncloned MRKAd5nef	MOI (vp/cells)
#1	B20010115-1	Uncloned	280
	B20010115-2	Uncloned	280
#2	B20010202-1	Cloned	280
	B20010202-2	Cloned	280

Results - Table 22 and 23 show an the ability to scale up production of MRKAd5nef by growth in a bioreactor.

Table 22: Virus Concentration as measured by the AEX assay 15

Run	Batch ID	Cloned/Uncloned	V	Virus Concentration @ 48hpi (1x10 ¹³ vp/L)						
		MRKAd5nef	Supernatant	Clarified Lysate	Total	Triton Lysate				
#1	B20010115-1	Uncloned	0.72	3.26	3.98	5.76				
	B20010115-2	Uncloned	0.38	1.67	2.05	2.46				
#2	B20010202-1	Cloned	0.80	6.00	6.80	8.88				
	B20010202-2	Cloned	0.50	6.00	6.50	8.47				

Table 23: Virus Titers as measured by the QPA assay

Run	Batch ID	Cloned/Uncloned		Virus Concent	ration @ 48hpi	(1x10" TU/L)	
		MRKAd5nef	Whole Broth	Supernatant	Clarified Lysate	Total	Triton Lysate
#1	B20010115-1	Uncloned	0.13	1.12	1.76	2.88	11.28
	B20010115-2	Uncloned	0.14	0.73	1.54	2.27	5.86
#2	B20010202-1	Cloned	0.14	0.97	1.62	2.69	11.89
	B20010202-2	Cloned	0.14	1.17	1.70	2.97	12.47

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The present invention is not to be limited in scope by the specific embodiments described herein. Indeed, various modifications of the invention in addition to those described herein will become apparent to those skilled in the art

from the foregoing description. Such modifications are intended to fall within the scope of the appended claims.

EXAMPLE 28

MRKAd5HIV-1gag Boosting of DNA-Primed Animals

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Groups of 3-5 rhesus macaques were immunized with (a) 5 mgs of V1Jns-Flgag (pVIJnsCMV(no intron)-FL-gag-bGHpA), (b) 5 mgs of V1Jns-Flgag formulated with 45 mgs of a non-ionic block copolymer CRL1005, or (c) 5 mgs of V1Jns-Flgag formulated with 7.5 mgs of CRL1005 and 0.6 mM benzalkonium chloride at weeks 0, 4, and 8. All animals received a single dose of 10e7 viral particles (vp) of the MRKAd5HIV-1gag at week 26. Note: 10e7 is too low to prime or boost effectively when used as a single modality (dose is selected to mimic preexposure to adenovirus); see Figure 32.

Blood samples were collected from all animals at several time points and peripheral blood mononuclear cells (PBMCs) were prepared using standard Ficoll method. The PBMCs were counted and analyzed for gamma-interferon secretion using the ELISpot assay (Table 24). For each monkey, the PBMCs were incubated overnight either in the absence (medium) or presence of a pool (called "gag H") of 50 20-aa long peptides that encompass the entire HIV-1 gag sequence.

The results indicate that MRKAd5HIV-1gag was very effective in boosting the T cell immune responses in these monkeys. At week 28 or 2 weeks after the viral boost, the number of gag-specific T cells per million PBMCs increased 2-48 fold compared to the levels observed at week 24 or 2 weeks prior to the boost.

The PBMCs were also analyzed by intracellular gamma-interferon staining prior to (at week 10) and after the MRKAd5gag boost (at week 30). The results for select animals are shown on Figure 31. The results indicate that (a) immunization with DNA/adjuvant formulation elicited T cell responses which can either be balanced, CD4⁺-biased or CD8⁺-biased, and (b) boosting with the MRKAd5gag construct produced in all cases a strongly CD8⁺-biased response. These results suggest that boosting with MRKAd5HIV-1gag construct is able to improve the levels of antigen-specific CD8⁺ T cells.

Table 24. Boosting of DNA/Adjuvant-Primed Rhesus Monkeys with MRKAd5gag

Number of SFChrilling PBMCs

Grey

Grey

T=0,4,8 wks DNA5 mgs PBS (D101)						,			51	1	÷L	t	Νl.		٩L			
DNA5 mgs PBS (D101)	T=26 wks		Medium	Dan H	Medium	GAG H	Medium	E E	medium edium	딁	틹		Ē	D89 H	E		Medium	
PBS (D101)	MRKAd5gag(E3+)	CBSH	NA A	AN	3	ક્ષ	5	7	4	224	-	116	-	8	_	330	0	316
(0101)	10.7 vp	X	_	0	•	5	•	46	0	8	•	92	-	33		1705	_	755
		AW3G	40	=	•	8	69	51	60	48	8	69		89	_	889	•	388
The second secon									_					1				
DNA/5mgs +	MRKAd5gag(E3+)	ည်	0	4	-	8	0	111	9	270	P	280	8	222	6	959	9	1345
CRL1005/45mgs	10.7 vo	8 ¥	4	0	-	101	0	284	٥	791		482	•	321	0	1915	-	<u>68</u>
		AW3P	6	60	-	9	4	7	4	164	_	ğ	9	æ	=	838	9	241
		CBSF	Š	×	0	9	0	88	0	530	19	374	6	251	В	1549	8	173
		AKBB	ø	12	4	8	_	119	•	439		425	0	316	4	1229	- -	1354
										7	1		1	1			1	
DNA/5 mgs+	MFKAd5gag(E3+)	AWZO	9	4	_	28	ທ	284	9	425	9	<u>\$</u>	60	202	6	565	8	\$
CRL 1005/7.5 mas + 0.6 mM BAK	10^7 vo	CARR	-	0	- -	121	-	135	_	270	9	뚕	_	105	=	<u>\$</u>	2	978
		CBSB		g	•	9		119	0	27.4	9	282	-	208	•	638	-	858
		CRSW	4	(*)		8	-	91	.0	139	•	20	_	8		643	_	349
		CB7D		. 0	•	136	•	316		60		929	_	769	•	2278	-	<u>5</u>
nane	Nome	960201	E	0	0	0	-	0	0	0	0	-	-	2	3	0	0	0

EXAMPLE 29

Construction of gagpol fusion for MRKAd5gagpol fusion constructs

The open reading frames for the codon-optimized HIV-1 gag gene was fused directly to the open reading frame of the IA pol gene (consisting of RT, RNAseH and integrase domains) by stepwise PCR. Because the gene (SEQ ID NO: 38) does not include the protease gene and the frameshift sequence, it encodes a single polypeptide of the combined size of p55, RT, RNAse H and integrase (1350 amino acids; SEQ ID NO: 39).

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The fragment that extends from the BstEII site within the gag gene to the last non-stop codon was ligated via PCR to a fragment that extends from the start codon of the IApol to a unique BamHI site. This fragment was digested with BstEII and BamHI. Construction of gag-IApol fusion was achieved via three-fragment ligation involving the PstI-BstEII gag digestion fragment, the BstEII/BamHI digested PCR product and long PstI/BamHI V1R-FLpol backbone fragment.

The MRKAd5-gagpol adenovirus vector was constructed using the BglII fragment of the V1R-gagpol containing the entire ORF of gag-IApol fusion gene.

EXAMPLE 30

Immunogenicity Studies in Non-Human Primates

Cohorts of three (3) macaques were immunized with 10e8 or 10e10 viral particles (vp) of one of the following MRKAd5 HIV-1 vaccines: (1) MRKAd5gag; (2) MRKAd5pol; (3) MRKAd5nef; (4) a mixture containing equal amounts of MRKAd5gag, MRKAd5pol, and MRKAd5nef, or (5) a mixture of equal amounts of MRKAd5gagpol and MRKAd5nef. The vaccines were administered at weeks 0 and 4.

The T cell responses against each of the HIV-1 antigens were assayed by IFN-gamma ELISpot assay using pools of 20-aa peptides that encompass the entire protein sequence of each antigen. The results (Table 25) are expressed as the number of spot-forming cells (sfc) per million peripheral blood mononuclear cells (PBMC) that respond to each of the peptide pools.

Results indicate the following observations: (1) each of the single gene constructs (MRKAd5gag, MRKAd5pol, or MRKAd5nef) is able to elicit high levels of antigen-specific T cells in monkeys; (2) the single-gene MRKAd5 constructs can be mixed as a multi-cocktail formulation capable of eliciting very broad T cell responses against gag, pol, and nef; (3) the MRKAd5 vector expressing the fusion

protein of gag plus IA pol is capable of inducing strong T cell responses to both gag and pol.

Table 25. Evaluation of Mixtures of MRKAd5 vectors expressing humanized

HIV-1 gag, pol, gagpol, nef in rhesus macaques

Grp#	gag, pol, gagpol, nef in rhesus mac	Monk#			T=6 wks		
	T=0, 4 wks	1	Mock	Gag H	Pol - 1	Pol - 2	Nef
1	MRKAd5 gag	CB9V	0	15		-	•
· 1	10^10 vp	CD19	0.	374	-	-	-
,	·	109H	1	843	-	-	•
2	MRKAd5 gag	99D130	1	948	•	•	-
1	10^8 vp	W277	16	324	-	•	-
		143H	4	595	-	-	•
3	MRKAd5 pol	CC1X	4	•	46	256	_
_	10^10 vp	AW3W	3		463	550	-
	·	AV43	6		95	1333	-
4	MRKAd5 pol	AW38	1	 	19	30	-
	10^8 vp	CC8K	0	-	50	995	-
		CC21	1	-	33	436	-
5	MRKAd5 nef	076Q	9	 -	-	 	1204
-	10^10 vp	091Q	4	-	-		85
:		083Q	0	-	-		176
6	MRKAd5 nef	000029	1		-	-	114
	10^8 vp	98D022	6	•	-		170
		98D160	3	-	-	-	198
7	MRKAd5gag+MRKAd5pol+MRKAd5nef	99D251	3	206	15	193	120
	10^10 vp each	05H	3	135	21	9	638
		00C016	3	26	4	51	23
8	MRKAd5gag+MRKAd5pol+MRKAd5nef	99D215	1	171	18	193	240
	10^8 vp each	81H	5	73	6	14	243
		12H	8	1140	115	811	719
9	MRKAd5gagpol +MRKAd5 nef	99D211	0	83	56	838	725
	10^10 vp each	22H	4	385	119	1194	1915
		61H	4	343	11	765	853
10	MRKAd5gagpol +MRKAd5 nef	34H	3	78	19	5	75
	10^8 vp each	48H	1	65	105	46	43
		70H	5	158	15	220	191

Indicated are numbers of spot-forming cells per million PBMCS against the peptide pools. Mock, no peptides; gag H, fifty 20-aa peptides encompassing p55 sequence; pol-1, 20-aa peptides representing N-terminal half of IA pol; pol-2, 20-aa peptides representing the carboxy-terminal half of IA pol; nef, 20-aa peptides encompassing the entire wild-type nef sequence. Responses to the antigens prior to the first immunization did not exceed 40 sfc/10⁶ PBMC.

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WHAT IS CLAIMED IS

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A recombinant adenoviral vaccine vector at least partially deleted in
 E1 and devoid of E1 activity, comprising:

- a) an adenovirus cis-acting packaging region corresponding to from about base pair 1 to between from about base pair 400 to about base pair 458 of a wildtype adenovirus genome; and
- b) a gene encoding an HIV protein or immunologically relevant modification thereof.
- 2. A vector in accordance with claim 1 comprising a packaging region corresponding to from about base pair 1 to about base pair 450 of a wildtype adenovirus genome.
- 3. A vector in accordance with claim 1 further comprising nucleotides corresponding to between from about base pair 3511 to about 3524 to about base pair 5798 of a wildtype adenovirus genome.
 - 4. A vector in accordance with claim 3 comprising base pairs corresponding to 1-450 and 3511-5798 of a wildtype adenovirus genome.
- 5. A vector in accordance with claim 4 which is deleted of base pairs451-3510.
 - 6. A vector in accordance with claim 1 which is at least partially deleted in E3.
 - 7. A vector in accordance with claim 6 wherein the E3 deleted region is from base pairs 28,133-30,818.

8. A vector in accordance with claim 1 wherein the gene encoding the HIV protein or modification thereof comprises codons optimized for expression in a human.

- 9. A vector in accordance with claim 1 wherein the vector comprises a5 gene expression cassette comprising:
 - a) a nucleic acid encoding a protein;
 - b) a heterologous promoter operatively linked to the nucleic acid encoding the protein; and
 - (c) a transcription termination sequence.
- 10. A vector in accordance with claim 9 wherein the gene expression cassette is inserted into the E1 region.
 - 11. An adenoviral vector in accordance with claim 9 wherein the gene expression cassette is in an E1 parallel orientation
- 12. An adenoviral vector in accordance with claim 9 wherein the gene expression cassette is in an E1 antiparallel orientation.
 - 13. An adenoviral vector in accordance with claim 9 wherein the promoter is a cytomegalovirus promoter devoid of intronic sequences.
 - 14. An adenoviral vector in accordance with claim 13 wherein the promoter is an immediate early human cytomegalovirus promoter.
- 20 15. An adenoviral vector in accordance with claim 9 wherein the promoter is a murine cytomegalovirus promoter.
 - 16. An adenoviral vector in accordance with claim 9 wherein the transcription termination sequence is a bovine growth hormone polyadenylation and transcription termination sequence.

17. An adenoviral vector in accordance with claim 9 wherein the transcription termination sequence is a synthetic polyadenylation signal (SPA).

- 18. A cell comprising the adenoviral vector of claim 1.
- 19. Recombinant, replication-defective adenovirus particles harvested and purified subsequent to transfection of the adenoviral vector of claim 1 into a cell line which expresses adenovirus E1 protein at complementing levels.

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- 20. An HIV vaccine composition comprising purified adenovirus particles of claim 19.
- 21. An HIV vaccine composition of claim 20 which comprises aphysiologically acceptable carrier.
 - 22. A method of producing recombinant, replication defective adenovirus particles containing the adenoviral genome of the adenoviral vector of claim 1 which comprises introducing the adenoviral vector into a host cell which expresses adenoviral E1 protein, and harvesting the resultant recombinant, replication-defective adenovirus.
 - 23. A method according to claim 22 wherein the cell is a PER.C6® cell.
 - 24. A method of generating a cellular-mediated immune response against HIV in an individual comprising administering to the individual a vaccine of claim 21.
 - 25. A method according to claim 24 which further comprises administration to the individual a DNA plasmid vaccine, optionally administered with a biologically effective adjuvant, protein or other agent capable of increasing the immune response.

26. A method according to claim 25 wherein the DNA plasmid vaccine is administered to the individual prior to administration of an adenovirus vaccine.

- 27. A method according to claim 24 wherein the adenovirus vaccine is
 5 preceded by an adenovirus vaccine of a different serotype.
 - 28. A method according to claim 24 which comprises administering and readministering the adenovirus vaccine vector to the individual.
 - 29. An adenoviral vector in accordance with claim 1 wherein the HIV protein is HIV gag or an immunologically relevant modification thereof.
- 30. An adenoviral vector in accordance with claim 9 wherein the gene expression cassette comprises an open reading frame encoding an HIV gag protein or immunologically relevant modification thereof.
 - 31. A recombinant adenoviral vaccine vector at least partially deleted in E1 and devoid of E1 activity, comprising:
 - a) an adenovirus cis-acting packaging region corresponding to from about base pair 1 to about base pair 450 and from about 3511 to about 5798 of a wildtype adenovirus genome, and deleted for base pairs corresponding to from about base pair 451 to from about base pair 3510 of a wildtype adenovirus genome; and
 - b) a gene expression cassette comprising
 - i) SEQ ID NO: 29;
 - ii) a heterologous promoter operatively linked to i); and
 - iii) a transcription termination sequence.

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32. An adenoviral vector in accordance with claim 31 wherein the gene expression cassette is in an E1 parallel orientation.

- 33 An adenoviral vector in accordance with claim 31 wherein the gene expression cassette is in an E1 antiparallel orientation.
- 34. An adenoviral vector in accordance with claim 31 wherein the promoter is a cytomegalovirus promoter devoid of intronic sequences.

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- 35. An adenoviral vector in accordance with claim 31 wherein the transcription termination sequence is a bovine growth hormone polyadenylation and transcription termination sequence.
- 36. An adenoviral vector in accordance with claim 31 which is at least partially deleted in E3.
 - 37. A cell comprising the adenoviral vector of claim 30.
 - 38. Recombinant, replication-defective adenovirus particles harvested and purified subsequent to transfection of the adenoviral vector of claim 30 into a cell line which expresses adenovirus E1 protein at complementing levels.
 - 39. An HIV vaccine composition comprising purified adenovirus particles of claim 38.
 - 40. An HIV vaccine composition of claim 39 which comprises a physiologically acceptable carrier.
- 41. A method of producing recombinant, replication defective adenovirus particles containing the adenoviral genome of the adenoviral vector of claim 30 which comprises introducing the adenoviral vector into a host cell which expresses adenoviral E1 protein, and harvesting the resultant recombinant, replication-defective adenovirus.

42. A method according to claim 41 wherein the cell is a PER.C6[®] cell.

43. A method of generating a cellular-mediated immune response against HIV in an individual comprising administering to the individual a vaccine of claim 21.

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- 44. A method according to claim 43 which further comprises administration to the individual a DNA plasmid vaccine, optionally administered with a biologically effective adjuvant, protein or other agent capable of increasing the immune response.
- 45. A method according to claim 44 wherein the DNA plasmid vaccine is administered to the individual prior to administration of an adenovirus vaccine.
 - 46. A method according to claim 43 wherein the adenovirus vaccine is preceded by an adenovirus vaccine of a different serotype.
 - 47. A method according to claim 43 which comprises administering and readministering the adenovirus vaccine vector to the individual.
 - 48. An adenoviral vector in accordance with claim 1 wherein the HIV protein is HIV pol or an immunologically relevant modification thereof.
- 49. An adenoviral vector in accordance with claim 9 wherein the gene
 20 expression cassette comprises an open reading frame encoding an HIV pol protein or
 immunologically relevant modification thereof.
 - 50. A recombinant adenoviral vaccine vector at least partially deleted in E1 and devoid of E1 activity, comprising:

a) an adenovirus cis-acting packaging region corresponding to from about base pair 1 to about base pair 450 and from about 3511 to about 5798 of a wildtype adenovirus genome, and deleted for base pairs corresponding to from about base pair 451 to from about base pair 3510 of a wildtype adenovirus genome; and

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- b) a gene expression cassette comprising
 - i) a nucleotide sequence selected the group consisting of SEQ ID NO: 1, SEQ ID NO: 5 and SEQ ID NO: 7;
 - ii) a heterologous promoter operatively linked to i); and

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- iii) a transcription termination sequence.
- 51. An adenoviral vector in accordance with claim 50 wherein the gene expression cassette is in an E1 parallel orientation.
- 52. An adenoviral vector in accordance with claim 50 wherein the gene expression cassette is in an E1 antiparallel orientation.

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- 53. An adenoviral vector in accordance with claim 50 wherein the promoter is a cytomegalovirus promoter devoid of intronic sequences.
- 54. An adenoviral vector in accordance with claim 50 wherein the transcription termination sequence is a bovine growth hormone polyadenylation and transcription termination sequence.

- 55. An adenoviral vector in accordance with claim 50 which is at least partially deleted in E3.
 - 56. A cell comprising the adenoviral vector of claim 49.

57. Recombinant, replication-defective adenovirus particles harvested and purified subsequent to transfection of the adenoviral vector of claim 49 into a cell line which expresses adenovirus E1 protein at complementing levels.

58. An HIV vaccine composition comprising purified adenovirus particles of claim 57.

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- 59. An HIV vaccine composition of claim 58 which comprises a physiologically acceptable carrier.
- 60. A method of producing recombinant, replication defective adenovirus particles containing the adenoviral genome of the adenoviral vector of claim 49 which comprises introducing the adenoviral vector into a host cell which expresses adenoviral E1 protein, and harvesting the resultant recombinant, replication-defective adenovirus.
- 61. A method according to claim 60 wherein the cell is a PER.C6® cell.
- 62. A method of generating a cellular-mediated immune response against HIV in an individual comprising administering to the individual a vaccine of claim 59.
 - 63. A method according to claim 62 which further comprises administration to the individual a DNA plasmid vaccine, optionally administered with a biologically effective adjuvant, protein or other agent capable of increasing the immune response.
 - 64. A method according to claim 63 wherein the DNA plasmid vaccine is administered to the individual prior to administration of an adenovirus vaccine.

65. A method according to claim 62 wherein the adenovirus vaccine is preceded by an adenovirus vaccine of a different serotype.

- 66. A method according to claim 62 which comprises administering and readministering the adenovirus vaccine vector to the individual.
- 67. An adenoviral vector in accordance with claim 1 wherein the HIV protein is HIV nef or an immunologically relevant modification thereof.

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- 68. An adenoviral vector in accordance with claim 9 wherein the gene expression cassette comprises an open reading frame encoding an HIV nef protein or immunologically relevant modification thereof.
- 69. A recombinant adenoviral vaccine vector at least partially deleted in E1 and devoid of E1 activity, comprising:
 - a) an adenovirus cis-acting packaging region corresponding to from about base pair 1 to about base pair 450 and from about 3511 to about 5798 of a wildtype adenovirus genome, and deleted for base pairs corresponding to from about base pair 451 to from about base pair 3510 of a wildtype adenovirus genome; and
 - b) a gene expression cassette comprising
 - i) a nucleotide sequence selected the group consisting of SEQ ID NO: 9, SEQ ID NO: 11, SEQ ID NO: 13 and SEQ ID NO: 15;
 - ii) a heterologous promoter operatively linked to i); and
 - iii) a transcription termination sequence.
 - 70. An adenoviral vector in accordance with claim 69 wherein the gene expression cassette is in an E1 parallel orientation.

71. An adenoviral vector in accordance with claim 69 wherein the gene expression cassette is in an E1 antiparallel orientation.

72. An adenoviral vector in accordance with claim 69 wherein the promoter is a cytomegalovirus promoter devoid of intronic sequences.

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- 73. An adenoviral vector in accordance with claim 69 wherein the transcription termination sequence is a bovine growth hormone polyadenylation and transcription termination sequence.
- 74. An adenoviral vector in accordance with claim 69 which is at least partially deleted in E3.
 - 75. A cell comprising the adenoviral vector of claim 68.
- 76. Recombinant, replication-defective adenovirus particles harvested and purified subsequent to transfection of the adenoviral vector of claim 68 into a cell line which expresses adenovirus E1 protein at complementing levels.
- 77. An HIV vaccine composition comprising purified adenovirus particles of claim 76.
 - 78. An HIV vaccine composition of claim 77 which comprises a physiologically acceptable carrier.
 - 79. A method of producing recombinant, replication defective adenovirus particles containing the adenoviral genome of the adenoviral vector of claim 68 which comprises introducing the adenoviral vector into a host cell which expresses adenoviral E1 protein, and harvesting the resultant recombinant, replication-defective adenovirus.
 - 80. A method according to claim 79 wherein the cell is a PER.C6[®] cell.

81. A method of generating a cellular-mediated immune response against HTV in an individual comprising administering to the individual a vaccine of claim 78.

- 82. A method according to claim 81 which further comprises

 5 administration to the individual a DNA plasmid vaccine, optionally administered with
 a biologically effective adjuvant, protein or other agent capable of increasing the
 immune response.
 - 83. A method according to claim 82 wherein the DNA plasmid vaccine is administered to the individual prior to administration of an adenovirus vaccine.

- 84. A method according to claim 81 wherein the adenovirus vaccine is preceded by an adenovirus vaccine of a different serotype.
- 85. A method according to claim 81 which comprises administering and readministering the adenovirus vaccine vector to the individual.
- recombinant, replication-defective adenovirus particles, wherein the adenovirus particles are harvested and purified from a cell line expressing adenovirus E1 protein, and wherein the particles are harvested subsequent to transfection of the cells with an adenoviral vector or vectors in accordance with claim 9; said vector(s) comprising a gene expression cassette or cassettes comprising nucleotide sequences encoding HIV proteins selected from the group consisting of:
 - a) gag, pol, and nef, expressed independently from three individual vectors;

 gag, pol, and nef, expressed independently from one vector with the encoding nucleic acid sequences operatively linked to distinct promoters and transcription termination sequences;

- gag, pol, and nef, expressed via two vectors, one expressing a polnef fusion, and another expressing gag;
- d) gag, pol, and nef, expressed via two vectors, one expressing a gagpol fusion and another expressing nef;
- e) gag, pol and nef, expressed via two vectors, one expressing a nefgag fusion and another expressing pol;
- f) gag, pol, and nef, expressed via one vector expressing a gag-polnef fusion;
- g) gag and pol, expressed independently from two individual vectors;
- h) gag and pol, expressed independently from one vector with the encoding nucleic acid sequences operatively linked to distinct promoters and transcription termination sequences;
- i) pol and nef, expressed independently from two individual vectors;
- j) pol and nef, expressed independently from one vector with the encoding nucleic acid sequences operatively linked to distinct promoters and transcription termination sequences;
- k) nef and gag, expressed independently from two individual vectors;
- nef and gag, expressed independently from one vector with the encoding nucleic acid sequences operatively linked to distinct promoters and transcription termination sequences;
- m) gag and pol, expressed via one vector expressing a gag-pol fusion;

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n) pol and nef, expressed via one vector expressing a pol-nef fusion; and

- o) nef and gag, expressed via one vector expressing a nef-gag fusion.
- 87. A multivalent adenovirus vaccine composition in accordance with claim 86 wherein the gag-pol fusion consists of SEQ ID NO: 39.
 - 88. A multivalent adenovirus vaccine composition in accordance with claim 86 wherein the fused sequences have the encoding nucleic acid sequences operatively linked to distinct promoters and transcription termination sequences.
- 89. A multivalent adenovirus vaccine composition in accordance with

 10 claim 86 wherein the fused sequences have the encoding nucleic acid sequences

 operatively linked to a single promoter; and the encoding nucleic acid sequences

 operatively linked by an internal ribosome entry sequence ("IRES").

Original Adenovector Construct:

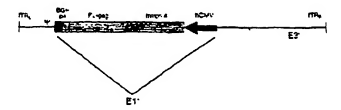


Figure 1: Original HIV-1 gag adenovector.

Sequence of the open reading frame for FL-gag (human codon optimized)

atgggtgctagggcttctgtgctgtctggtggtgagctggacaagtgggagaagatcaggctgaggcctggtgg caagaagaagtacaagctaaagcacattgtgtgggcctccagggagctggagaggtttgctgtgaaccctggc agctgaggtccctgtacaacacagtggctaccctgtactgtgtgcaccagaagattgatgtgaaggacaccaag gaggccctggagaagattgaggaggagcagaacaagtccaagaagaaggcccagcaggctgctgctggc acaggcaactccagccaggigtcccagaactaccccattgigcagaacctccagggccagatggtgcaccag gecatetececeggaecetgaatgeetgggtgaaggtggtggaggagaaggeetteteeetgaggtgateee catgiticitigecetigicigagggigecaeeeeeeaggaeetgaacacatgetgaacacagtgggggggccate aggetgecatgeagatgetgaaggagaceateaatgaggaggetgetgagtgggacaggetgeateetgtge acgciggccccattgcccccggccagatgagggagcccaggggctctgacattgctggcaccacctccaccct ccaggagcagattggctggatgaccaaccaccccccctctgtgggggaaatctacaagaggtggatcat ccigggccigaacaagatig:gaggatgtactcccccacctccatcciggacatcaggcagggccccaaggag cccticagggactatgtggacaggttctacaagaccctgagggctgagcaggcctcccaggaggtgaagaact ggatgacagagaccctgctggigcagaatgccaaccctgactgcaagaccatcctgaaggccctgggccctg gctgaggccatgtcccaggtgaccaactccgccaccatcatgatgcagaggggcaacttcaggaaccagag gaagacagtgaagtgcttcaactgtggcaaggtgggccacattgccaagaactgtagggcccccaggaaga agggctgctggaagtgtggcaaggaggccaccagatgaaggactgcaatgagaggcaaggccaacttcctg ggcaaaatctggccctcccacaagggcaggcctggcaacttcctccagtccaggcctgagcccacagcccct agetglaceceeiggeeteeetgaggteeetgtttggeaaegaeeeeteeteeagtaaaataaageeegggea gat (SEQ ID NO: 29)

Figure 2

Old Transgene:



New Transgenes:

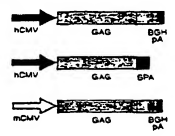


Figure 3: Diagrammatic representation of the original HIV-1 gag transgene and the series of new transgene constructions.

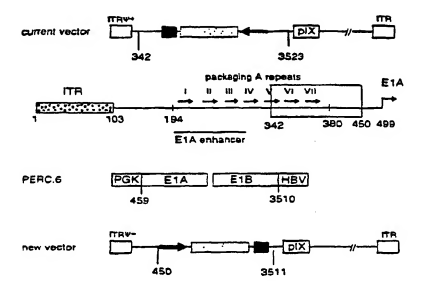


Figure 4: Modifications made to the current adenovector backbone in the generation of the new vector.

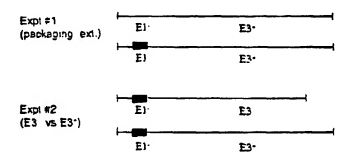


Figure 5: Virus mixing experiments to determine the effects of the addition made to the packaging signal region (Expt #1) and analysis of the effects of the E3 gene on viral growth (Expt. #2). The red bars denote the region of modifications made to the E1 deletion.

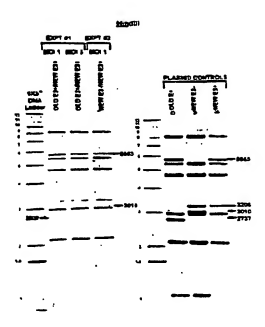


Figure 6: Autoradiograph of viral DNA analysis following viral mixing experiments (expts. #1 and #2) as detailed in the text.

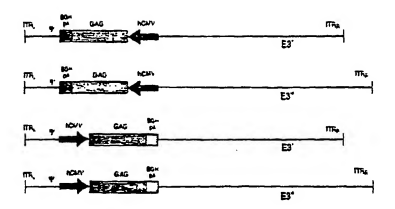


Figure 7A: hCMV-FLgag-bGHpA adenovectors constructed within the "MRK" backbone. E1 parallel and E1 antiparallel transgene orientation within the E3- and E3+ backbones were constructed.

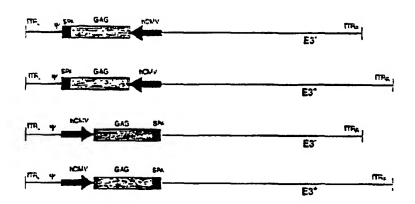


Figure 7B: hCMV-FLgag-SPA adenovectors constructed within the "MRK" backbone. E1 parallel and E1 antiparallel transgene orientation within the E3- and E3+ backbones were constructed.

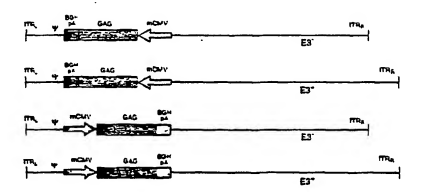


Figure 7C: mCMV-FLgag-bGHpA adenovectors constructed within the *MRK* backbone. E1 parallel and E1 antiparallel transgene orientation within the E3- and E3+ backbones were constructed.

Plasmid mixing expt: (orientation)

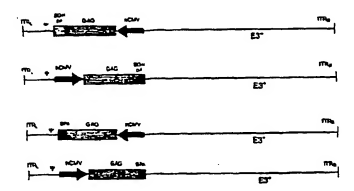


Figure 8A: Effect of transgene orientation

Plasmid Mixing expt: (poly A signal)

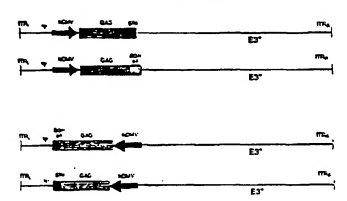


Figure 8B: Effect of polyadenylation signal

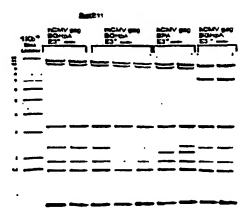


Figure 9: Viral DNA from the four Adgag candidates at P5, following BsfE11 digestion.

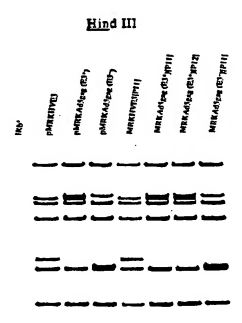


Figure 10: Viral DNA analysis of passage 11 and/or 12 of MRKHVE3, MRKAd5gag and MRKAd5gag(E3-).

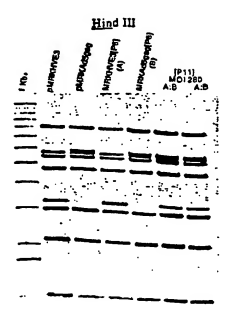


Figure 11: Viral DNA analysis (*Hin*dIII digestion) of passage 6 MRKHVE3 and MRKAd5gag used to initiate the viral competition study. Last two lanes are passage 11 analysis of duplicate passages of the competition study (each virus at MOI 280 vp).

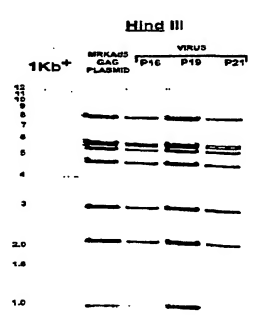
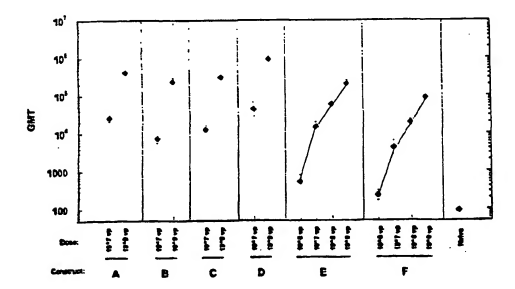


Figure 12: Viral DNA analysis by *Hin*dIII digestion on high passage numbers for MRKAd5gag in serum containing media with collections made at specified times. The first lane shows the 1 Kb DNA size marker. The other lanes represent pre-plasmid control (digested with Pac1 and *Hin*dIII), and MRKAd5gag virus continually passaged to P16, P19 and P21(serum containing media).

Figure . Serum anti-p24 Levels at 3 Wks post i.m. immunization of balb'c mice (n=10) with Varying Doses of Several Adgag constructs: (A) MRKAd5gag (through passage 5): (B) MRKAd5 E3* hCMV-FLgag-bGHpA; (C) MRKAd5 E3* hCMV-FLgag-SPA; (D) MRKAd5 E3* mCMV-FLgag-bGHpA; (D) research Lot (293 cell-derived) of Ad5HIV-1gag; and (F) clinical lot (Ad5gagFN0001) of Ad5HIV-1gag. Reported are the geometric mean titers (GMT) for each cohort.



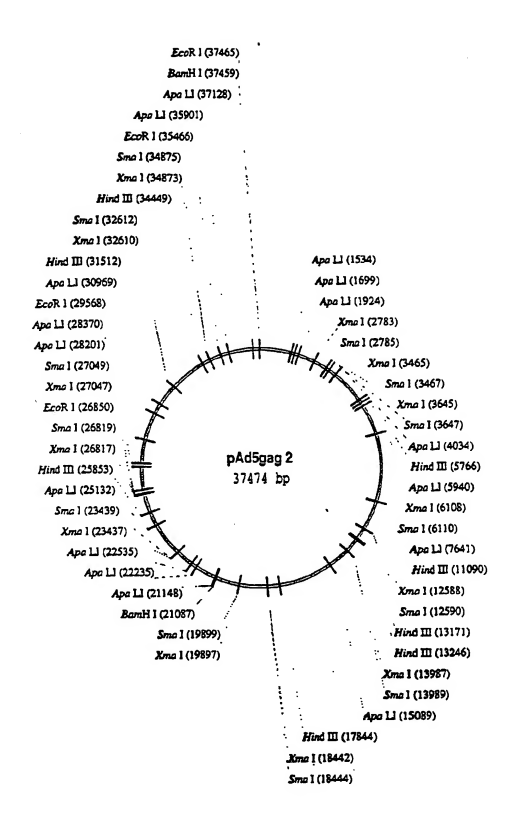


Figure 14

GCFCCCACCTC AAACACTTCA CCCCTCCCCC ACCCCACTCTCT TOTAL TOTALCTTT TCCCTCCCT ACCCTTTTT TCCCTCCCT ACCCCTTTTC ACCCCAAAAA CCCCTTTTTC ACCCCAAAAA CCCCTTTTTC ACCCCAAAAA CCCCTTTTTC ACCCCAAAAA CCCCTTTTTC ACCCCAAAAAA CCCCTTTTTC ACCCCAAAAAA CCCCTTTTTCC ACCCCAAAAAA CCCCTTTTTCC ACCCCATTTTCC	ICC COCATTORIC CATTOTANC CONTINUANCE CICCUTTING CT ACCCCCCCC GOACTTINAC COTTINOCIO DISACTICIA C INA PECCOCCCC CETGANICIO GEANITGEA CICTIANACE INA COCCCCCCC ATCCATICCA TACGTICIAT CCATATIAT INT COCCCCCCC TAGGTANCET ATCCATATA GATATATAT INT COCCCCCCC CAGANICAT ATCCATATATI	ICT CATCITATTA ATACTAATCA ATTACGGGGT CATTACTICA ICT CATCAATAAT TATCATTACT TAATGCCCCA GTAATCAAGT ICC GCCCAACGAC CCCCCCCCAT TGACGTCAAT AATGACGTAT ICG CGCCAACGAC GIGGCGGGTA ACTGCAGTTA TAACAGCAAA	CHI TANACTIVICCE ALTTGGCAGT ACATCAAGTG TATCATATY* ALE ATTTGACTGG TGMCGGTCA TOTAGTTCAC ATAGTATACT WIT ACATGACCTT ATGGGACTTT CCTACTTGGC ACTACATCTA ICA TGATCATGGAA TACCCTGANA GGATGAACCG TCATGTAGAT	ATARCECCANA CTENERGOO GATTECCANO TATCECCANA CTENERGECC CTANAGETTC AACANCTECE ECCENTIGAE GENANTGGE TITITITAGEE GOODTANCTO COTITACEE	TIG CGCTACCACG CTGTTTGAC CTCCATAGAA GACACCGGTAA TIG CGGTAAGTGC GACAAAACTG GAGGTATCTT CTGTGGCCCTT Bigili TAGAATCTAC CATGGGTGCT AGGCTTCTG TGCTGTTTGG TAGAATCTAC CATGGGTGCT AGGCTTCTG TGCTGTTTGG	THE CONTINUED ATTOROGOGY CCTCCARGON GUTTINAMING THE CONTINUED TAKENGARCE CONGENERAL CONCULTIVE TAKENGARCE CONCULTIVE CONCULTATION ACCITECACIO CONTINUED
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ACATCATCAA TGTAGTAGTT GTAGTAGTCT CATCATCACA GGAAGTGACA	CCACATGIGI CCTTCACTGT TAAAACTYAAACTYAAACAAAAAAAAAAAAAAAAAAAA	TTATATIGGC ANTATANCCG ATGGAGTTCC TACCTCANGG	TAACGCCAAT ATTGCGGTTA CCCTATTGAC GGGATAACTG		TACCOTICOGA GOTCTATATA AGCAGAGETE ATOCCACCET CEAGATATAT TEGITETEGGG CEGATECAGE CTECTOCOGCE GOGAACAGTG CONTACTAGE CACCAGAGATATAGAGAGAGAGAGAGAGAGAGAGAGAGAG	GACAGTGGG GACAGTGGG ACCCTGGCC TGGGACCGGA CACAGTGGCT GTGTCACCGA AAGGCCCAGC
101 C C C C C C C C C C C C C C C C C C	301 401 401	501 A T T 109	701 G 801 G	901 C	1101 . T	1301 T 1401 T 1501 C

Figure ISA

DNRKAI'IGAG HERGB2

1701	CACCAGGCCA	TCTCCCCCC	GACCCTCAAT	CCCTCTCTCA	Actorization	ראזאהאאהפרכ	THETECTOR	ACCITCATCCC	CATOTTCTCT	GCCCTGTCTC
1801	ACCONCOCAC	CCCCCARGAC	CTCAACACTA	TCC TCAACACAC	ATATATA SECUL	CATACTUCACTE COTACT COT	CCATCCACAT CCTACCTCTA	GOTTONGOOD	ACCATCAATG TOGTAGTTAC	ARABARGETER TECTTECTAL
1901	TGAGTGGGAC	AGGCTGCATC TCCGACGTAG	CTCHYICACKS	TOXCUCCATT	ממנגמנוטטטט	ACATY:ACCCA	CATCCACCACAC	TCTGACATTG	CTOCCACCAC	CTCCACCC
2001	CAGGAGCAGA	TICCCTCCAT	GACKINALING		CHTTHYSOCKT	ANTETACANG	ACCACCTAGE	TCCTOSCCT	GANCAAGATT	GTCAGGATTE
2101	ACTCCCCCAC TOAGGGGGTG	CTCCATCCTG	GACATEMORIC		הבאהכהרדה הבחנומאאה	AGGACTATG	TGGACAGGTT	CTACAAGACC	CTCAGGGCTG	AGCANGCCT
2201	CCAGGAGGTG	AAGAACTOGA	TCACAGAGAC	CCTACTAGTG	CAGAATGCCA	ACCCTGACTG	CANGACCATC	CTGAAGGCCC	TOGRECCTOC	FOCCACICETY ACCENTAGON
2301	CTCCTCTACT	TOACAGCCTO	CCACCACACAC	GCCCCCAGAC	CHTACAAGGC	CAGGGTGCTG	GCTGAGAGCCA	TOTCCCAGGT	DACCAACTEC	GCCACCATC, CGCTGCTAG
2401	TGATGCAGAG ACTACGTCTC	GOGCAACTTC	ADGIAACCAGA TECTTGGTCT	GGNAGACAGT	CTTCACGAAG	AACTGTGCCA	AGGTGGGCCA TCCACCCAGT	CATTECCARD	AACTUTAGGG TTGACATCCC	CCCCCTARGIA.
2501	GAAGGGCTGC	TOGANGTOTO	GCANGGARGG CGTTCCTCCC	CCACCAGATG	MAGGACTACA	ATGAGAGGCA TACTCTCCGT	CCGGTTGMG	CTCCCCAAAA	TCTGGCCCTC AGACCGGGAG	CCACAACICA: GOTOTTCCCC
2601	AGGCCTGOCA TCCGGACCGT	ACTICCTCCA TGAAGGAGGT	GTCCAGGCCT CAGGTCCGGA	GAGCCCACAG	CCCCTCCCGA	CCTCAGGANG	ACCIANCECE	ACCACTACAG	CACCCCAGC CHUNGCAN	CACIANGEAR?
2701	AGCCCATTGA	CAAGGAGCTG	TACCCCTVIG	CCTCCTGAG	GTCCCTGTTT	CRICAAMTAACC CKISHTKK.HGG	CCTCCTCCCA	GTANMTAAA CATITITATIT	OCCCOGRCAG CGGGCCCGTC	ATCTOCTOTY: TAGACGACA
2801	CCTTCTAGT	GCCAGCCATC	TOTTOTTOC	CCCTCCCCCGGGCCCCCCGGCCCCCCCCCCCCCCCCCCC	TOCCTTCCTT	GACCTGGAA	GGTGCCACTC CCACGGTGAG	CCACTGTCCT	TICCTAATAA	ANTCACCOAN TTACTCCTTT
2901	TTOCATCOCA	TTGTCTGAGT	AGGTGTCATT TCCACAGTAA Pvul	CTATTCTVAG	CCCACCCCAC	RECORDACA CCCGTCCTGT	GCANGOTATA CGTTCCCCCT	GCATTOOGAA CCTAACCCTT	GACAATAGCA CIGITATCGI	GC:ATCCTCC CCCTACGACC
3001	GGATGCGGTG	GCCACTATOG	CCCATCGGCG	CCGATCGGG GGGCATGAGGGGGGGGATGAGGGGGGGGGG	AAATGIKITKG TTTACACACC	CACATCACTFA	AGGETGGGAA	AGAATATATA TCTTATATAT	AGCTGGGGGT TCCACCCCCA	CTTATOTAGE
3101	THGTATCIO ÁAACATAGAC	TTTTGCAGCA AAAACGTCGT	COCCOCCACC	CCATGAGGAG GGTACTCGTG	CAACTUGITT	CATGOAAACA CTACCTTCGT	THEM AND THE A	ATATTTCACA TATAAACTGT	ACOCCCATCC TGCGCGTACG	CCCCATGGGC GCCCTACCC+1
3201	CCCCCTCCCCT	CCCCCTCCCT CACAATCTCA	ACCURACE	CATTICATEMENT	CCCCCCCCTATC	TOTACCIDE AAA	CTCTACTACC	TRACCTACG	AGACCGTGTC TCTGGCACAG	TCCAACGCCG ACCTTGCGGC

tique 158

PMRKAdSgag MER682

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1101	THERESTALIS	THEORY CACICALORS	CANCELLER	STATEMENT STATEM	נבעניניניני	COSCAPICA	Activitation	CHITCHOAG	CCCACTIGCA	AACAC:TCCAG
	MCCTCTGAC	GICGGNGGCG	DCCCCCAACT	CCCHTACTTE: CCCTCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC	כא: בטאנאא		-			TIGHTCACGTC
3401	GAMERIALANG	ATCCCCCCC	CATCACAGE	ACTIVICACIONED ACTIVICACIONED	AAACCCICIT	AACCTMGAA	TCACCCCTACA ACTUACACCCT	ACTTANTOTO TGANTTACAG	CAAAGAGTCG	ACCINCITIVATA TOCACAAM "F
3501	TCTCCCCAG		CCCTV:AAGGC		CCCAATTACG	TTTANACAT		CCACACTCTG	TTREATTIE	GATCAAGCAA
1006	MACGCGGIC		Science Period	אייוייייייייייייייייייייייייייייייייייי	ACCORDERATE.	WILLIAM STATE			TAPPETER	ACCACCACCAT
3001	CACAGAACGA	CACAMTAMA	TCCCCANANC	GCTSCGCTCCA	TCCAGCCCT	COMPLETE			ATAMANAGE	TCCTGCACCA
							F _S (I)			
3701	AAAGGTGACT	GACCTACAAG	AGATACATCC TCTATGTACC	CCATANCICC	CAGACACCIC	THE AGGTAGE ACCTECATEG	NCCACTFICAG TOCTCACGTC	ACCTICATEC	TCCCCCCTCC ACCCCCCACC	ACAACATUTA
3801	GATCCAGTCG		acrosseera		Anchemenca	GTAGCAAGCT	GATTGCCAGG	GCCAGGCCCT	TCOTCTAACT	GTTTACAAAC
	CTAOCICAGE		CGACUTACAC	CACGGATTIT	TACAMANGT	CATCGTTCGA	CTAACGGTCC	CCCTCCCGGGA	ACCACATTCA	CANNIGITAE
3901	COCHTANCT	COCATOGGTO	CATACGTOOG	CATATGAGAT	GCATCTIVESA	CTUTATITIE	ACCITICACITA			CORRESATIVA
	OCCANTICGA	CCCTACCCAC	OTATICCACCC	CTATACTICTA	CGTWINACCT	GACATAAAA	TCCAACCGAT	ACANGOGICG		OCCCTANIT
4001	TOTTOTOCAG	-	ACAGITGITATIC	COGRECACTE	GGGAATTTG	TCATGTAGCT			_	COCCCTTCITY
	ACAACACOTC	Troorogred	TETCACATAG	CCCACCTGAA	CCCTTTANAC	AGTACATOGA	Archicelta	ACCCACCTTC	TICAACCICT	GCCCCAACAC
4101	ACCTCCAAGA	_	ATTCGTCCAT	AATGATGGGA	ATTATACTOR	ათაითიით				GTCATAGETO
	TOGAGGITCT	AAAAGGTACG	TANGCAGGTA	TTACTACCGT	TACCOCKITG	تحدوددوردو	-			CAGTATCAAC
4201	TOTTCCAGGA		ATAXXCATT	TTTACAAAGC	GCCCCCCCCAAG	ממדמככאנזאב				GCOTAGTTA
	ACAAGGTCCT	ACTETAGEAG	TATCCCKGTAA	MATISTITICS	מסכניטינית	CCACCGTCTG	ACCCCATATT ,	ACCAMGGTAG (C(X:ATC:AA'n:
4301	CCTCACAGAT	TRUCATT	CACCETTINGA		COCCATCATG					GCCIACIATE A
	OGAGICICIA	AACGTAAAGG	GTGCGAAACT	CAAGTCTACC	CCCCTAGTAC	AGATGGACCE	CCCCCTACIT	CTTTTGCCAA	AGGCCCATC	CCCTCTAGRIC
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	CHOCOMON	TCGTCCAAGG		GCTGAATGGC GTGGGCCACC	CHUCOCCACC		GTGTGGATAA			TICTCTCGAC
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4501	CAGCTOCCGT	CAGCTOCCOT CATCCCTOAG	CAGGARGCC	ACTICGITIAN GCATGICCCT	GCATGTCCCT	GALTCOLATG	TTTTCCCTGA	CCANATICCOC (TERECERCEA
	GTCCACOCCA	GTAGGGACTC	שבנבנבכבפ	TCANGCANTT CGTACAGAGA	CGTACAGGA	CTGAGGGTAC	GACT	GCTTTAGGCG (מוכדולכפכה	ACCOCCOOT
4601	OCCURTACCAG			THICMCDG	PTTGAGAGCG				CCANGCAGTT	CCAGGGGGTC
	COCTATOOTC	AAGAACGTTC	CFFCGFFFCA	ANANGERICACE	AAACTCTCASC	MERCENTA				A.J.C.C.C.
4701	CCACACCTCO		CTACGGCATC	TCRATCCAGE	ATATCTCCTC	CANACACCOCC	Progradence /	AMOCGACAT (CCACCAGTAGT (GCCACGAGGA
1007	GUTUTCOAGE		SATOCCOMO		Windle No.		_	_		כסריוניהכני
4801	GCTCTGCCCG	GICCCAGIAC	AGAMAGTAC		CCACC: ACTICG	CATCAGACCC		_		GCGACCGGT"

figure 15c

PMRKAdSgag MER682.

4901	CCACGCCTTC	ACCORDERACE	TOCTOOTICET ACCIACCACCA	GAAGERATIVE	CPGTCTTCGC	CHINCHINGTO	CCCCACTTAG	GYTTAGACTAGE	ACCACACTAT	CACCACACACA
2001	TCCGCGGGGT			_	AGRICAL EACH	CTINGUESTICAGE (A TYCCCGTC	TOCACACTIT	THACHCCCCTA	CAGCTTVACC	CCTAGAAATA
5101	CCGATTCCCG				GARGEAGAGE	CATTCCACGA	GCCACTAG	CHCTGGCCGT	TCCGGGTCAA	AMACCAGGIT
5201	TCCCCCATGC AGGGGGTACG	STITITGATGE MANACTACG	GTTTCTTACC	TCTCATTTCC	ATVANCOCCE	GTCCACGCTC	GCTCACGAAA	AGGCTGTCCG	TOTCCCCTA TACACHCTY ACAGGGCAT ATGTCTGAN	TACACACETY: ATGICTGAM:
5301	AGAGGCCTGF TCTCCGGACA	-	TGTTCCRCGG ACANGGCCCC	TCCTCCTCGT	ATAGAMACTC TATCTTTGAG	GAACCACTCT	CACACAAAGC	CYCGCUTCCA	OCCCAGCACO	AACKIACICTA
5401	AGTOGGAGGG TCACCCTCCC	OTACCOCACC CATCCCCACC	TROTCCACTA	CCCCAGGTG	TCGCTCCAGG ACCAMOGICC	CACACTTCTB	ACATETCOCC TOTACAGCOG	CICITICOCCA	TCAAGGAAGO	TOATTOSTIT
5501	CATCCACATC	OCCACOTOAC COGTOCACTO		TGAAGGGGGG	CTATAAAACG	GACTORGGGC	CCCAMICAGE	TCACTCTCTT	CCCCATCCCT	GTCTCCCACC
5601	GCCAGCTGTT	•		- •	TCACTTCTGC	GCTAAGATTB	TCAGTTTCCA AGTCAAAGGT Himilii	AAAACGAGGA	DOATTIGATA CCTAAACTAT	PPCACC-FGF7
5701	CCGCGGTGAF GGCGCCACTA	OCCITITIONOO COGNANCICC	Gracecert caccaceta	CCATICTOSTIC GGTAGACCAG	MGMAMAGACA TCTTTTCTGT Pvol	ATCTTTTTGT TAGAMARCA	TOTCAACCTT ACAGTTCGAA	CCACCOTTE	GACCCGTAGA	GENERATION CCCCCAACCT
5801	CACCAACTTO	GCGATGGAGC	CCACCCAAAC	CANAMICACO	CCATCACCC	GCTCCTTGGC	COCCATOTT	ACCTOCACOT	ATTCCCCCC	AACGCACCCA:
5901	CATTCOOGAA	AGACOOTGOT	OCCUPATION CO	GOCACCAGGT	CCACCCCCCA	ACCICCOCTTO	TOCAGGGTGA	CANCOTCAAC	OCTOOTGOCT CGACCACCGA	ACCTCTCCG TCCAGAGGC
6001	OTACOCOCTC CATCCCCCGAG	GITGOTCCAG	CAGAGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGG	COCCCTITATO	CCACCAGAAT	GACGATAGAG	GGICTAGGIG	CONCINCTICC	CCCCCCAGAC	CCTCCACCACT
6101	MAGACCCC	OCCACCAGOC CCOTCOTCCO	GCGCGTCGAA	GTASTCTATC	THECATECTT	GCAAGTCTAG	CCCCTCCTGC	CATOCOCOGO	COCCANDECE	CCCCTCGTA1
6201	CCCAACTCAC	CCCCTGGGGT	TOCCATOCOO ACCGTACCCC	TRACTITACED ACCENCING	CGGACGTA	CATCCCCCAA	ATCTCCTANA	CCTAGAGGGG	CTCTCTGAGT	ATTCCAAGA? TAAGGTTCTA
6301	ATGTAGGGTA TACATCCCAT	OCATICITICSA COTAGAAGOT	CCCCCCANTIC	TRACIACIAC AC ACCACACAC	GTAATCGTAT	ACTTCGTGCG TCAAGCACGC	MASSAGECING TECETEGETE	OAGOTCOGGA CTCCAGCCCT	CCGAGOTTOC	TALIGGGCCCC
6401	CHOCHCHOCH	CCCANCACTA	TCTCCCTCAA AGACGCACTT	GATGACATGT	CHEANCETAC	ATATOCTIVES	ACCCHICANG TOCCACCTTC	ACCTTOMACE TOCAACTTCG	TRANCOTICHOT ACCCCAGACA	DAGACTTACK: CTCTCX:ATC7:

VICA PACT ICT : SGA :	LTV':	וובני מפפה מכני	GCGr trict: reag neag	SACC	2000	בפאת פכדי	NCTC FGAG	CTCA	CATY	: כבוכ		LLV.
TCCTTGATGA AGGACTACT CGTCGGCCT GCAGCCGAG			CANGAAGCG" OFFCFIX:FC1: CCAAGATFAG COFFCFACTIC:			AGCACGCT		CITCGICTCA	ACCENCATY: TRECACCEN: TRECTEGONG		AAGGTCCCCG	
GICCAGAGITT CAGGTCCCAA ATCCAAAACC TAGCCTTTGG	GGACGCGCCG	•	TOTAAAGTTC ACATTTCAAG GOCCCAGTCT CCGGGTCAGA			AAAGAGACGC 1711CTCTGCG	GRADANGTEC CTGCGACGGG CATETICAGG GACGETGCEC	ACCTGACGAC CGCGCACANO TGGACTGCTG GCGCGTGTTC Khol	CTRACTOCTC GARGOGAGTT GACCGACGAG CTCCCCTCAA GACCACAGATAA GACTGTCCA		CTATGGATTA	
CCCCCCCCCCTAT CCCCCCCCTCAT CTACTCTTCC CATCAGARACC	TCGCGCATAC CGCATCCGCC	GCGTAGGCGG AGGCATAAAG TCCGTATTTC	TCCCCACAA ACCGCCTCTT GCTCTGAAAG CGAGACTTTC		OCCOCAOTCA CCCCGTCAGT	CCTARGTGAC					AGATCCAGGT	האיניהואה
Treaggreta Acgretare GGTCTTTCA CCAGANAGOT	AAGATCACCA TCAGTGTCGT	ACTUACACACCA TTY CUCCUCO AACKACACCOCOC	GTTGATGTTG CAACTACAAC CTCAGCCCGT GACTCGGCA		TAGGITCTCGC ATCCAGAGCG	GTCTCTACAT		CACGAGGTTG	CCTTROACCGT CXTAACTKIGGA	ACTICATION	GACKRETCAG RACGCRAGCT CTRECCAGTC CCUCCCCGA Knnl	מאידארייהייא הדאהיהיה הכליהיה הכליהיים
	TEXTTAGGGAA GTATTTGAAG	CATAAACTTC AAGAGTATCT TTCTCATAGA	CCTCAAACCC GCACITTCGG TTCACCCGAG		CCANACCCC	CCANGTATAG GGTTCATATC		GTACATCCTG CATGTAGGAC			GACKAGTCAG CTCCCCAGTC KnM	מאיידאניאניויא
•	TCCTAGGGGGG ACCATCCCTT TGAGATACTT		AGCACGATCT TCGTCCTAGA AGGTCAGCTC		TCCCATCCAA AGGGTAGATT	AGGCCCCCATA TCCCCCCCCTA	ATTOCACTOR FAACCTCCTC	TGCACTACCT ACGTCCCC()A		CHECKEGGGA	ACCTCOCATA TOCACCCTAT	נכבטכטיטינונ
	GTTGACGACCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC	-	CTCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC		Trecendera Anggotece	TCCTTCCCAA ACGAAOGGTT	CCCGCCACCA	CTGGCAGCGG GACCGTCGCC	-	AGATGTCCCC TCTACAGGCG Psil	CTGCAGGTTT GACGTCCAAA	COCTTOCANG AGGICCICATC CICCYCCHARDIC
GTACKIAGTEG CATCETEAGE TITTITITEC AAAAAAAGG	TGTAGAACTG ACATCTTGAC GGTGTCCCTG		TOTTANTHAC ACANTTANTG CAATTITITA	-	OCCIONATO PARA COCCIONACIONAL COCCIONACIONAL COCCIONAL C	CCCCACCACC	AACTGGATCT	GACGCGACTA		CCCAAAGTCC	_	_
CCANGGAGGC GCTTCCTCCG ATCCTGTCCC TAGGACAGGG	GAGCCTAGCA CICCGATCGT TGAGCGCAAA	ACTCGCGTTT TTTGGAACGC AAACCTTGCG	TCGGAACGGT ACCTTGCCA TGATGGAAGG	CHOCHANCE	TACAAGOTAA	CCAGCATGAA GOTCOTACTT		CHOCHGECT THOTAAAAC		GCCGCGCGAG	CTCCCGCGC OTCAGGTCAG GAGGGCGCCG CAGTCCAGTC	TREPTRESTUS COCCUTCGAT
	CCAACOGTAA CCTTOCCATT GAGGTOTOGG	CTCCACACCC CCGNGCGCTT GGCACGCGAA	TCCCGGCACC AGGGCCGTGG GGGATGCCCT	GOTTOGNAGC CCAACCTTCG	CCACTACOTC	AACTTCATOA TTOAAGTACT	GATGCGAGCC CTACGCTCGG	GROCTGGCTT	CCCTTAACT	CCTOGIGGIO	CTCCCGCGGC	TOCHROSTOR
6501	6701 6801	6901	7001	7201	7301	7401	7501	7601	1701	7801	7901	8001

PMRKAd5gag MER682

8101	ATCCATCTAA	Anococycac	GCGCGCGAGC	CCCCCCCCACACT	AGRETATION	בננאפעככנינכ	CCCCAGACCC	GOCAGGGGGA	CONCERNICACE	GCGCGCGCFAC
8201	ASSASCTUST				בינאנדאיסטים מרשביטרונפי		THANFORME		GAAGACGACG	CCCCCTTCA
8301	GCTTGAACCT		-	CAATTTCCCT		מינייינידאיני	GCTTTTAGAG		CCTGAGTTGT	CTTCMTAGG: GAACTATCC:
,										
8401	CTAGAGCCC	ATGAACTOCT	CCATCICITIC	CHUCHNEAGA	TCTCCCCCTC	COSCINGENCIA	CACOGINGES	CCCACCTCCCT	TCGRAATCCG ACCTTTACGC	CCCATTACTO:
8501	TOCCHOAGO				TETTAGACCAC	GUCCUCATICG	GCATCGCATG		CACCTIGCOCO	ACATTOACT:
8601	CCACGTOCCO	GCCGAAGAC			AAAGAGGTAG	THEACACHES	TCCCCCACAC		AAGAAGTACA	TANCCCACC!
8701	TCCCAACGTG	GATTCOFTG))	OCCUTCANGG	CGCTCCATOR	CCTCGFAGAA	GTCCACGGCG		ACTORGAGET	GCCCCCCGAC
	AGCOTTGCAC	CTANGCAACT	ATAGGGGGTT	CCGGAGTTCC	OCCACACTACC	GRAGCATCTT	CAGGTGCCGC	TICANCTER	TOACCCTCAA	בפכנונים
8801	ACCONTANCT	CCTCCTCCAG	AAGACGGATG TTCTGCCTAC	AGCTCGGCGA TCGAGCCGCT	CAGTOTCOCO	CACCTOROROGO	TCANAGGCTA AGTTTCCGAT	CAGGGGCCTC	TTCTTCTTCT	TCAATCTCE:
6									A CONTOCKACO	, A
1068	CHECCATANG	CCGGAGGGGA	AGANGANGAN	GACCOCCOCC	ACCCCCTCC	CCCTCTCCCG	בכשבתשכתשב	במכטבספבבב .		GTTKGRAA
9001	GATCATCTCC	CCCCCCCCAC	CCCCCTACCA	CAGCCACTGC	GTGCGGGCGT CGCGCCTTCA	Tritciacogna AGNACOCCC	GCCCAGTICAGC	AAGACGCCGC TTCTGCGGCG (CCOTCATGTC	CCCASTTATION
9101	GTTOOCOOOO CAACCGCCCC		CGCCAGGAT	ACCOCCOCTAA	CGATGCATCT	CACAATTOT	TOTOTAGGTA ACACATCCAT	CACCOCCOCC	GACCCATC	AGCGAGTCT** TCGCTCAGG
9201	CATCUACCOG	CATCGACCOG ATCGGAAAC GTAGCTGGCC TAGCCTTTTG	CTCTCGARMA GAGAGCTCTT	ACCCCAGATT	CCAGTCACAG	TCCCAARGTA	G3CTGAGCAC CCGACTCGTG	CGTGGCGGGC (GCACCGCCG)	CCTCGCCCG	GGCGCTCCC#: CCGCCAGCCC
9301	CAACAAAAAC	GCGGAGGTGC	TGCTGATGAT	GTANTTAMG	TACKRESSIFICE ATEXXXCAGA	TOAGACGGGG ACTCTGCCGC	CTACCAGCTG	AGAAGCACCA TOTTCOTTCOTTCOTTCOTTCOTTCOTTCOTTCOTTCOT	TOTACCTIFICED ACAGGAACCC	TCCGGCCTCC
9401	TOANTECOCA ACTTACOCOT	OCCOCTCOC	CAMPCECCAG	CGANGCANAN	GACATCHINCE	CACCAGAAAC	TACTACTOR	GCATGAGCT COTACTCGGA A	TTCTACCOOC AAGATOCCCO	ACTTCTTCT TGAAGAAGA
9501	CTCCTTCCTC	ACAGGACGE	TCTCTTGCAT AGAGAACGTA	CTATCRCTOC GATACKC	בנגאלנעכנטכ בנגאלנעכנטכ	CTCAAACTAG	CATCCACCC	CCCTCTTCCT CCCCCCCCCCCCCCCCCCCCCCCCCCC	CCCATGCGTG	TGACCCCGA.A ACTGGGGGCT F
9601	OCCCCTCATC COOCAGTAG	GOCTGAAGCA CCGACTTCGT	GRACTAGOTC CCCGATCCAG	CCCCTOTTICC	CCC.TraditA CCCACCCCAT	ATATCCCCCAC TATACCCCCAC	CPCCACCTOC GACGROSPICE	GICAGACITNG I	ACTURANDTC TGACCTTCAG	APCCATCHCY TAGGTACAGG

9701	ACAAAGCGGT	GETATOCGCC	ACAAAGCOGT GGTATOCGCC CGTGTTGATG GTGTAAATCK		ACTRICICCAT		TTAACGGCTCT	GETTACCCOS CTOCHAGAGC	CTOCCIAGAGC	TCGSIGIAL C.	
	TOTTTCGCCA	CCATACGCG	G GCACAACTAC	CACATTICACO	TCAACCCAITA	TRICCTOTIC	AATTGCCAGA	ANTIGICAGA CCACTORGCC GACRETCTCG ACCCACATAS	GACGCTCTCG	ACCCACATUS:	
		Xhol	lor mare								
9801	TGAGACGCGA	GTANGCCCT	C GAGTCAAATA	CCTFACTICATT		ACCIVICATION GOTATCICCAC		CAAAAAGTGC GGCOGCGGCT		COCOLITACAY:	
	ACTOTOCOCT	CATTCGGGAG	CTCAGTTTAT	GCATCACTA	COLLECTOR	TEXTRICATEIA CCATAGGGTG	CCATAGGGTG	GITTITICACG	CCCCCCCCGA	CCCCATCTC	
				Refil		Feafi	_}				
9901	GGCCAGCGT	AGGGTGGCCG	OCCUCADO		TECAACATAA		TCCCTACIATE		TCCAGGTGAT	מכנפפנטונפ	
	CCCCOTCCCA	TCCCACCOGC	CCCCAAGGCCC	CCCCTCTAGA	AGGITGIATT	CCCCTACTAT	AGSCATCTAC	ATCGACCTGT	AGGTCCACTA	دوددوددود	
10001	CTRACTROCAGE	CCCCCCCCAAA	GTCGCGGACG	CCCTTCCAGA	TOTTRECTION	CCACAAAAAG	TGCTCCATTGG	TCGGGACGCT	CTOOCCOOLC	Arteriorecto.	
)))	CACCACCTCC	GCCCCCT		GCCAAGGTCT	ACAACGCGTC	GCCGTTTTTC	ACCINCITACE	ACCCCTGCGA	GACCGGCCAG	المحدود بحدود :	
		Xbal									
10101	AATCOTTOAC	8	GTOCANAGG	AGAGCCTGTA	AGCORPAGE	CTRECORGE	CTCATAGATA	AATTCCCAAG	COTATCATOR	COCACGACCC	
	TTAGCAACTG		CACOTITICE	TCTCGGACAT	TCGCCCGFCA	CHACACACLA	GACCACCTAT	TTAAGCGTTC	CCATAGTACC	eccuecates.	
10201	COUNTROADE	CCCOTATECG	OCCUPACE DE	GTGATCCATG	COUNTACCIO	CCCCCTRTCG	ANTCCARTIG	TREGARGICA	GACAACGGGG	CAGTGCTCC-T	
	CCCAAGCTCG		COCCAGGCCAG	CACTAGGTAC	GCCANTRACIO	GACCACACACC	TRACETICCAC	ACCCTOCAGE	CHCHIOCCCC	CTCACGAGGA	
10101	THOCTICE		OCCOCCIOCT	GCGCTAGCTT	THITKARCAC	Transcored	CMACGIANGE	GGTTAGGCTG	GANACICGANA	GCATTANGTY:	
	MACCOMOG			CGCGATCGAA	MANACCOSTG	Arcagagaga	GICCCATICG	CCAATCCGAC	CTITICOCTIF	CGTAATTCAL	
10401	(STRUCTUCE)		OCCITATITI	CCAAGOSTTG	AGRICACTION	CCCCCGGTTC	CACTCTCCCA	CCIGCCGGAC	TOCOGOGOANC	GGGGGTTTGF	
	COAGCOAGG			CONTRACTAR	TCACCCCCT	GOTTOCCANG	CTCAGAGCCT	OCCCOCCIO	ACCCCACTIG	CCCCCANCI	
10501	CTCCCCGTCA		CGCTTGCAAA	TTCCTCCGGA	AACAGGGACG	AGCCCCTTTT	TICCETTICC	CAGATOCATO	COSTOCTOCO	GCAGATGCG	
	GAGGGGCAGT			AAGGAGGCCT	TRINCCCTGC	TCGGGGANA	AACGAAAAGG	OTCTACGING	GCCACGACGC	COTCTACGC	
10801	مددنساداد			CAGCGGCAGA	CATECAGGGC	ACCTCCCCT	CCTCCTACCG	CCTCAGGAGG	OCCGACATCC	GCGGTTCACC	
	CCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC			GICCCCGTCT	GTACCTCCCG	TYXCARGURA	GGAGGATYKKC	GCAGTCCTCC	CCCCTGTAGG	CCCCACTYX	
10701	COSTAGEAGA			2000000000	CCCCCACTAC	CIRGACTING	AGGREGGEGA	0000010000	COOCTACOR	COCCUTATION	
	OCCUTORE	ACCACTAAT	_	ככשכשכככנ	COCCUTOATO	CACCTGAACC	Treverent	CCCCASACCOC	OCCONTACT	GCGGGGAGA(*;	
10801	TITACACACACAC		AGCTUANGCO	TOATACGCGT	CACCUCTACG	TCCCCCCCATA	GAACCTGTTT	COCCIACCOCCI	ACCCACACACA	CCCCCACACAC	
	ACTOCOCOTO			ACTATOCOCA	CICCGCATGC	ACCIOCOCCIOT	CTTCCCACANA	(acacTOOcoc	receneration	COGOCICCIC	
10001	ATTENDED		CGCAMGGGG	GAGCTGCCAAC	ATCCCTCAA	TECCEARCE	THICHGOOD		TONGCCCGAC	CCCCCAACCC	
10601	TACCCCCTAG			CTCCMCCCCC	TACCCCACTT	ACCICCICC	AACGACGCCC	TCCTCCTGAA	ACTOCOCOTO	S COCOCUTOGO	
										Section 1	
11001	GOATTAGTCC	CONTINUTCE COCOCOCCA	CACGTGGCGG	CCGCCGACCT	PCTAACCIPCA	TACGAGGAGA	CCCATCAACCA		TTTCAAAAA	THEMANA CETTINGAN	
	CCTAATCAGG	acacacacat	GTGCACCGCC	CACCOCTCRIA	CCATTGGGGF	ATCCTCCTCT	GCCACTICGT	CCICTANTIG	MANGELLILI		
111101	CCACCTICCGT	· ACCCTTOTOO	COCCCCANGA	GGTGGCTATA	GRACTGATIC		CTTTTTANGE	GCGCTGGAGC	MAMCCCAM	TNOCHARIS C.C.	
! !	GOTGCACGCA	TOCCAACACC	: occcenter	CCACCGATAT	CCTGACTAGO	TAGACACCCT	CANACATTCG	CCCCACCICO	THE COLUMN	HICKITICANI.	
11201	CTCATOGCGC		TATAGTGCAG	CACAGEAGE		ATTCACCCAT	GURTICITA	ACATACTAGA	GCCCGAGGGC	COCTOCCING	
	GAOTACCOCO	-	ATATCACGTC	GIUTCUICCE	TGITTGCTCCG	TAAGTCCCTA	CGCGACCATT	TOTATCATCT	נפשבורופ	Crowce war	

			******			a creation of the control of	1004000 W		THE STATE OF STREET	THE BELL PROPERTY.
11301	TCGATTIGAT	TCGATTIGAT AMCATCCTG	CAGAGLATAG	איזכראבראריה.	CH CH TAN THE	AL AL TABLE DE SE	COUNTY THE TAIL	CCCATAGENG	ATANGGTACG	AATCGGACC
	ACCTAMACTA	ACCTARACTA TITICTACCAC	GICILLIAIC	ACT. ACT. ACT.	16161111		4.7.4.4.4.4.4.4.4.4.4.4.4.4.4.4.4.4.4.4			
11401	CAACTITITAC	CCCCCCCAAGA	TATACCATAC	CCCTTACGITT	CULTAINIACA	ACCIPACITANA	OpenVariate	TTCTACATGG	CCATCCCCCT	CAMINITAL
	GTTCAMATO	COCCCTICT	ATATCCTATC	GOGNATOCAA	CATATATE IN:T	TTTA'SCALLE	CTACKTECKE	ANGANDTACO	CCTACCCCCA	CTTRICACCI .
11501	ACCITICAGEG	ACCACCTGGG	CCITITATICA	MCGMACGCA	TEXTAINANCE	כניוניאוניכוני	M XCCGGGGGGC	GCGAGCTCAG	CGACCCCCAAG	CTYSATTACACA
	TOOMACTOGO		GCAANTANCO	TRICTCGCGT	AGGICTRICG	GCAL: TCGC: AC	אימפכבמכנים	COCTCGAGTC	GCTGGCGCTC	GACTACOTAST
11601	GCCTGCAAAG	COCCCTGGCT	GGCALTRACA	GCGCCCATAG	NUNGCUCUYC	TCCTACTTTC	ACGCGGGCGC	TGACCTGCGC	TGGGCCCCAA	GCCCAACTCGC
1	COGACOTTIC		CCGTACCCGT	CACCIONTATO	וגיוכנפפנונ	ACCATGAAAC	TOCACCCACG	ACTGGACGCG	ACCCCCCCCTT	כמסכוסכטכ:
11701	CCTCGAGCA	OCTOOOGCCG	GACCTGGGCT	GACGGTAGCA	נברניניניניניניניניניני	CTGGCACGT	COCCACCOTO	GACCAATATO	ACGAGGACGA	TCACTACGAG
	COACCTCCCT		CTOGACCCCA	CCGCCACCGT	ממינטניטטינכ	GACCGTTGCA	OCCCCCCAC	CTCCTTATAC	recreeract	ACTCATGCTC
									- File	
11801	CCAGAGGACG	OCCAGIACTA	Accordanta	TITCTGATCA	CATCATGCAA	GALFICAACTS	ACCORDOR	0000000000	CTGCAGADCC	AGCCGTCCC:
	OGICTOCTIGO		TCGCCACTAC	AMGACTAGT	CTACTACGTT	CTGCGTTKKC	TREGRECOTATEA	coccaccac	GACGICTOGG	TCGGCAGGCF
11901	CCTTAACTCC	ACGGACGACT	GCCCCNOT	CATOGACCIAC	ATTATETECE	TRACTIFICATO	CAATCCTAAC	acontocoac	AGCAGCCGCA	GOCCANCON!
1	GGAATTGAGG		CCCCCCTCCA	GTACCTGGCG	TAGTACAGUG	ACTGACGCGC	GTTAGGMJTQ	COCANGOCCO	TCGTCGGCGT	COOCHIOCC.
							Tage			
12001	CICICISCAA	TICTICARGO	OCTOOLECCO	GCGCGCGCAA	ACCCCACCCA	CGAGAAGGTG	CTOCCCATUG	TAMACGCCT	GCCCGAAAAC	AGGCCATCT
	CAGAGGCGILT	ANGA	_	COCCOCOTT	reconteen	ACTOTACCAC	GACCICCTARIC	ATTTOCCCCA	CCCCCFTTTG	TCCCGGTAG
12101	CHECCERACGA	900000000	GTCTACGACG	CACTGCTTCA	CCCCCTCCCT	CCTTACAACA	GCGGCAACGT	GCNGACCANC	CTYXOACCOCC	TOTOGOGGA
1	CCCCCCCCCC		CAGATGCTGC	GCGACGANGT	CCCCCCCCC	CCANTITITIT	CCCCCTTCCA	CCICTOGTIG	GACCTGGCCG	ACCACCCCC"
12201	TOTOCOCOAG		AGCGTGAACG	CCCCCACCAG	CAGGGCAACC	TOCOCTOCAT	GGTTKACACTA	AACOCCTTCC	TGAGTACACA	OCCCOCCAN'
	ACACGCCCTC		TCCCACTCGC	accentate	GTCCCGTTGG	ACCCCANGGTA	CCANCGTONT	TTGCGGMGG	ACTCATGTOT	COCCOSTE
12101	OHOCOOCOO		CTACACCAAC	TTRUTCHOCG	CACTGCGACT	AATYBGTYGACT	מאלאכאכנפכ	AAACTGAGGT	GTACCAGTCT	COCCCAGACT
	CACOGCGCCC		CATOTICCTIO	AAACACTCGC	GTGACGCCTA	TTACCACTGA	CICICIONOCO	TITCACTCCA	CATOGICAGA	CCCCCTCTGA
			P.4	_						
12401	APPERECA	GACCAGTAGA	CAAGAGCCTGC	NGACCGTAAA	CCTGAGCCAG	ACTITICAMA	ACTICCARGO	CCTCTCCCCCG	GRECOGETE	CCACAGGCGA
	TAMANAGGT		GTTCCGGACG	TUTCHCATTF	GGACTCGTTC	CHANAGETETE	TRANCOTOCO	CCACACCCCC	CACGCCCCAG	CONCICCOCT
12501	CCGCGCGACC	GIGICIAOCT	TOCTGACACC	CAACTEGGGG	CTGTTTATTGC	TYCTAATAGE	GUTTERE	GACACTOGCA	OCOTGICCCG	GGACACATAC
	OCCOCCIO	CACAGATCGA	ACGACTGCGO	CTTCAGCGCG	מאכאאכמאכנ	ACGATTATCG	CHARIANGTER	CTCTCACCGT	CCCACAGGGC	CCTGTGTATA
12601	CTAGETICACT TEE	TEXTOACACT	GTACCGCGAG	GCCATAGATIC	AGGRECATET	CHALLANCIAT	NUTTICCARG	AGATTACAAG	TOTCAGCCGC	מכמכוניםמימכ
	CATCCAGTGA	ACCACTGTGA	CATCCCCCTC	COGTATCCAG	TECCHESTACA	CCTXXTTCGTFA	THINANGETEC	TCTAATOTTC	MCMGTCGGCG	COCCOVICCO
								•	Pireji	
12701	AGGAGGACAC	AGGAGACAC GOOCAGCCTG	CAGACANCEC	TAAACTACCT	CHITCHACTANG	CHACHACAGA	AGATOCOGTO	GTTCCACACT	TTAMACAGCG	ACCACCACC
	recreeters	recreations eccorcooke	CTCCOTTIX	ATTRIBATION	רכאכתיאידות	CULTATORICE	TUTACKARAG	CAACGTGTCA	MTTMTCGC	Techerna.
12801	CATTITIOCOC	CATITIOCUE PACOTOCAGE	AGAGCCTTCAG	CCTTAACCTG	ATTACORTS	CHATTAACINIC	CARCITICATIO	CTCCACATGA	הכתכסכת: א	CATOSAACTA
	GTANANCGCG	OTAMACOCO ATOCACOTOS	TCTVITCACTC	CRONATTORDAC	TACTACKTING	CCCATTGCGG	מוכנאניעניניני	מאנירדיקדאנד	GACACUTE	פוארני

12901	COCATOTATO	CCTCANACCR	OCCUPITATIC	AACCOCCTAA	TYXIACTT	מכעונומנאנט	Strict Control	ACCCCCMOTA	THEACCHAT	CCCATCTICA
	CCCTACATAC	GGAGTTTGGC	CCCCANATAG	TTKKKYKKYTT	ACC"PI:ATCAA	CCTACCCACA	こいいことのことのこ	TYSTACTOAT	MACHGOTTA	CYCTACIARCT
13001	ACCCRCACTO		CCTCATFICT	ACACCCTTTT	ATTRICACATE	COTCACERTA	ACGATCRATT	CENTROCAC	GACATAGACG	ACACACOTOTA
	TOCOCUTOAC	CGATCCCCCC	GGACCANAGA	TOTAL	TAAGCTCCAC	COSTITUTORY	TKICTACCTAA	CHAGACCCTG	CTOTATCTGC	TGTCCCACAA
								Hingail		
13101	THECECOGAA	CCGCAGACCC	TOCTAGAGAT	GCAACAGGG	GAGCAGGCAG	ARRICHICA:T	GCGAAAGGAA	GCGAAAGGAA AGCTTCCCCCA	GOCCHAGCAG	CTIGICCGAIL
	AAGGGGCGFF	AACOGGCGTT GOCGTCTGGG	ACCATCTCAA	CGITICACCCC	دعدوعددوعد	TUCKSCCGCCA	CUCTITICOT	CUCTITICCTT TCGAAGGCGT	CCGGTTCGTC	GAACAGGCT'.
					United	٠				
13201	CTAGGCGCTG	COCCCCCCC	GTCAGATGCT	AGTAGCCCAT	TTCCANGITT	GATAGGGGTCT	CTTACCAGCA	CTCCCACCAC	CCCCCCCCCC	CTCCTOTIGC:
	DATCCCCCGAC	DATCCGCGAC GCCGGGGCGC	CAGTCTACGA	TCATCCCCTA	AAGCTTCC:AA	CTATCCCAGA	GAATGGTCGT	GAGCOTGGTO	0000000000	CACCACCC
			PS-d	_**						
13301	ACCACCACTA	CCTANCANC	tegetherine	איכניניכאמנים	CGANANAMC	ניתככידכה	CATTTCCCAA	CAACGOGATA	GAGAGCCTAG	TOGACANIA1
	TCCTCCTCAT	COATHOTTO	AGCGACGACG	TOGGOGINGO	GCTTTTTTG	מעכהמאמניככ	GTAMAGGCT	GTTGCCCTAT	CTCTCGGATC	ACCTN:TTCTA
13401	GAGTADATUG	AAGACGTACG	CCCAGGAGCA	CACCCACCTO	CCAMACTICC	GUCCIACCAC	CCGTCGTCAA	AGCACGACC	GTCAGCGGG	rendendina
	CTCATCTACC	TICTOCATGO	OCGICCTICOT	GICCCTGCAC	GGTYTTCKARCG	CCARRECAGIO	GGCAGCAGTT	TCCGTGCTGG	CAGTCCCCC	AGACCACACT
13501	GAGGACGATG	ACTOGGCAGA	CCACAGCAGC	GPCCTGGATT	TROCHANGE	TYCONCCCG	TTRECTERE	TTCGCCCCAG	GCTCGGGGAGA	ATGTTTTAAA
	CTCCTGCTAC	TOAGCCGTCT	GCTNFTCGTCG	CAGGACCTAA	ACCCTCCC.TC	ACCOUNTGOOD	NANCTACGTUR	AAGCGGGGTC	COACCCCTCT	TACAMATIT
13601	MANAANAA	GCATGATGCA	ANATAANNAN	CTCACCAACK	CCATTAXCACC	GAGCCTTRGGT	TITICITYSTAT	TCCCCTTAGE	ATCCCCCCC	COCCEATGTA
	Trustana.	COTACTACGE	TTTATTTT	GAGTGGTTCC	GGTACCCCTCC	CINCCAACCA	ANAGAACATA	AGGGGAATCA	TACGCCGCGC	GCCGUTACAT
13701	TOAGGAAGGT	CONCORDE	CCTACCIACIO	TOTOGRAPH	המאממניתה	TRANCOCCAGO	CK:TYGGFTCT	CCCTTCGATG	CTCCCCTOGA	CCCCCCTTT
	ACTCCTTCCA		GGATCCTCTC	ACACCACTCG	COCCOCOGIC	ACCGCCCCCG	CCACCCAAGA	COGRACCIAC	GAGGGGGACCT	GOCCOCM.
		Kins								
13801	oraccreces		GCCTACCORG	CHCAGAAACA	CCATCCGTTA	CTCTGAGTTG	CCACCCCTAT	TEGACACEAE	CCGTOTGTAC	CTGGTGGACA
	CACCCACCCC	CCATGGACGC	COGATOCCCC	CCCTCTTIGE	CCTAGGCAAT	GATACTENAC	CCTCCCGGATA	ACCINENCINO	GCCACACATO	GACCACCTOT
13901	ACANGTCAAC		TCCCTTAACT	ACCAGAAGGA	CCACACCAAC	TTTCTCACCA	CCGTCATTCA	AAACAATGAC		GRGAGGCAAG
	TOPTCACTIC	CCTACACCGT	NOCCACTICA	TOSTCTICCT	actorcarta	AMAGACTECT	CCCAGTAAGT	THEITACTE	ATCICIOSOCC	CCC1CCC11.
14001	CACACAGACC	ATCANTCTTO	ACONCCORTC	GCACTRAGARC	CKICCACCTICA	AAACCATCCT	GCATACCAAC	ATCCCAAATC	TOANCOAGIT	CANTITIACE
	GIGTOTOTOG	TAGTTAGAAC	TGCTCGCCAG	CONCACCCCG	CCUCHAGACT	TTTKGTAGGA	CGTATCCTTC	TACOCITITAC	ACTITOCTICAA	GTACAMTEG
14101	ANTAAGTTIA	AGGCGCGGGT	GATOCTUTES	COCITICACTA	CTANGGACAA	TUNCTIVENG	CTCANATACG	ACTICOCTORA	OFTICACGCTG	CCCGARRACA
	TTATTCAAAT	TCCGCGCCCA	CTACCACAGC	OCCUNCOGAT	GATTCCTVTT	AGTCCAUTTC	GACTITATEC	TCACCCACCT	CAAGTGCGAC	GOGCTCCCGT
				ł	Pari			•		
14201	ACTACTOCOLA		ATAGACCTTA		GATCHINENG	CACTACTEGA	AACTGROCAG		_	CACCONCATICOS
	TOATCAGOCT	CTOGTACTOG	TATCTICAAT	ACTIVITIES	CTAGE:ACI:TC	CHEATGAACT	TTCACCCGTC		CARGACCTTT	CIRCHITAGOC
14301	OCTANACTIT		ACTTCAGACT	CONTITIONS	دندىيدىدى	GRETHINGAT	CCCTVAXXXTA		_	TCCAGACATU
	CCATTTCAAA	CTGTGGGCGF	TGAAGTCTGA	בכבראאוניהו	באניורטע־אנאנ	CAGNACAGTA	CORPUCCEAT	ATATCTTICC		MASTETATAG
14401	APPTROCTOC	CACCAT	AGINAGACTTC		מיב אימענאט		ATCITICATARGE			TTTAGGATCA
	TAMARCACG	GTCCTACGCC	CCACCTICIANG	TOXXITCHOC	COCACTOCATE	מאכאינכנג	TACKACCITICG	CCCTTCCCAA	CONCENCE OF	MATCCTAGT

Figure 15I

14501	CTTBOOKTOB	Transactories of	CCCPAACATTIC	CONTRACTOR	CATANTACAC	CHETAIRCANI	CGAGCTTGM	AGATCACACC	CANCAGGGCG	מספנונטינוני
10041	CCATGCTACT		CCATTOTANG				GCTCGAACTT	TCTACTGTGG	CTIVITCCCGC	CCCCACCGCT
14601	AGGCGCCAGC	AACAGCAGTG	GCACACACAC	ניניאעניאניאעני	TUCAACGETS:	ויאנאיריאנו	אאוניאנונכם	GTGGAGGACA	TRAACGATCA	TOCCATTON:
	TCCCCCCTCG	TIGICOTOR	CONCIOCUACO	CCTTCTCTTG	AGGTTROCCATC	(מולמויתאונים	TTACGINGGE	CACCTCCTGT	ACTTOCTAGE	ACCCTAACCC
14701	CACCORCACO		COCTGACGAG	AACACACCAT	אטטטנעטאטט	Marchitann	פון די אוניברא דיכ	CCCCTCCCC	ACCCOAGOTC	GAGAAGCCTA
	CCUCTOTOGA		CCCACTCCTC	TTCGCCCCAC	Teconomics	TCCCCGRCTT	CHACKICHIG	GOCGACGCGT	TGGGCTCCAG	CICTICGGW
										Krati
14801	AGANGAACC	COTCATCAAA	CCCCTGACAG	NATACAGGAA	CANACICAGE	TACAACCTAA	TAMECANTOA	CAGCACCTTC	ACCCAGTACC	CCAC-TC-T'A
	Perietrico	PETTICITIES CCACTAGITT			CTTTGCCTCA	ATCITICALITY	ATTECTTACT	GTCGTGGAAG	TOCOTCATOO	CGTCGACCAT
	K Da									•
14901	CCTTGCATAC	AACTACGGCG	ACCUTCAGAC	COGMITTEE	TEATGRACE	TYCTTINGAC	TOCTOACCTA	ACCTGCGGCT	COCHOCAGO	CTACTCAGTCC:
1	GGANCOTATO		TOCOACTOTO	accttaaca	AGTACICITIOG	ACTIMAACTITE	ACKSACTYKYAT	TOGACOCCOA	accreateca	GATCACCACC
10031	THEORY BUBER	TOATOTARDA	CCCCTICACC	TRUCCULOCA	このことにいいいと	CARCATT	CCGGNACTOG	GCCCCGAGCT	OTTOCCCGTO	CACTCCAAGA
	ACCOTOTO		-		GCGCGCTCTA	GTCGTTGAAA	GTCCACCACC	CCCGGCTCGA	CAACGGGCAC	OTCAGGTTCT
•										***************************************
15101	CATHETACAA	CONTRACTA CONCEAGOC	OPETACTECE	AACTCATOOG	CCAGTTTACC	TETETGACCE	ACCINOTICAL	TCCCTITCCC	GAGAACCAGA	THITCOCK!
70171	COANGATOTT	geroorecad			GGTCANATGG	AGAGACTGGG	TOCACAGGTT	AGCGAAAGGG	CICITOGICI	AAAACCGCC"
	Asci									
15201	CUCCAGC	CCCACCATCA	CCACCGTCAG	TGARANCOTT	CCIVICITICA	CAGATCACGG	CACCCTACCG	CTGCGCAACA	GCATCGGAGG	AGTECAGETIA
	GOOCGOTCOC	000100			GCACGAGAGT	GTCTAGTGCC	CTYCGATOGC	CACCCCTTCT	CCTAGCCTCC	TCACCTCCCT
16101	GROACCATTA		ACCCCCCACC	TGCCCCTACG	TTTACAAGGC	CCTCACACATA	GICICOCCOC	GCGTCCTATC	GAGCCCCCACT	TTTTGAGCAA
	CACTGGTAAT				MATCTTCCO	CHACCCCTAT	CAGAGCGGCG	COCACCIATAD	CTCGGCGTGA	AAAACTICITT
15401	Car American		_	ACACACACTG	שאגבונונפנ	TTCCCAAGCA	AGATGITIOG	COCCCCAAG	AAGCGCTCCO	ACCAACACO.
40,50	COTACAGGTA	CCAATA	_		CCCOGACGCG	AACCONTACCT	TCTACAAACC	OCCCCBGTTC	TTCGCGAGGC	recructor:
***					CACAAACGCG	GCCGCACTIG	מנמכעכנענב	GTCGATGACO	CCATCOACGC	CONTRACTOR !
10661	Transfer and			-	GROTTIGCGC		CCCCTCCTCC	CAGCTACTOC	GGTAGCTGC	CCACCACCT.
15501	Characters		_		CACTOCACGC	CCCCATTCAG	ACCOTOGIC	GCTAGAGCCCO	GCGCTATGCT	MANTGANGA
10001	CHECKOO		-		GTCACCTGCG	CCAGTANGTC	TRICACCACG	כפנכובסססכ	CCCCATACGA	THINCETCI
15701	GALCHICAGE		_	CCCCCCGACC	CHICACTRICC	מבכנאטנאנו	ටයාටරාවරයට ර	CCTCXCTTAAC	CACACATO	פכאכניאיניכני
	CTGCCGCCTC		GCAGCGGTGG	COCCOCTO	CICCCTGACGG	CAGGITICOCC	פבבעבעבעב	GCACCAATTG	GCGCGTGCAG	COMPCION
		Sfi								
15801	ACOGGCOGCC	ATGCGG	CTCGAAGGCT	· GACCGCGAOT	ATTRICTURE		GTCCAGCCGA	CCACCOCCCO		CACARCATI
•	100000000	TACCCCCCCC	GACCTITCCGA	י בכמפכשבבע	TAACAGINIAC	ACCACACAC	CARGITCCGCT	GUICECCAR		CACAC.CONT.
15901	AGTOCTATICA	CTCAGGGTCG	CAGGGGGAAC	GRETATTER	-		CTGCGCGTGC	CCGTGCGCAC	ددودددده	CUCAACTA
	TCACGATACT		GICCCCG1TG	CACATAMICC	ACCCCCTGAG		מאכפכיטכאכפ	GACACICUIS	GA WASHING	
16001	TTGCAAGAAA				TECARCACA		ACCANGCTAT	Checkwaeae	THERETTE	TICTCTACKA
	AACGITICITIT	F TITICATICAAT	CTGAGCATGA	CANCATACAT	AGGTCGCCGC	מאככנאינאכניד	והכוורהאוא	באיין וכמכנו		

Figure 15J

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	CCAGGTCATC	CCAGGTCATC GCGCCGGAGA	TCTATROCC	TCTATCOCC CCCGAAGAAG GAACAACAG ATTACAAGC CCGAAAGCTA AAGCGGGTCA	いいくしていているい	ATTACAACCC	CTCAAACCTA	ANGCOODICA	MAAGAMA	GNANGATICAT	
	GOTCCAGTAG	COCOGCCTCT	AGATACCGGG	OGGUTETTE	CTT TRIGING	TAATGITTCKS CKICTTTCGAT	CCTPTCGAT	TREGECEAGE	minchine	CTTTCTACT.	
								Sati			
16201	GATGATGAAC	TTOACCACCA	CONTROVICTO	CTCACCCTA	ניג גייטעניו, כיכי עני	הההארואיהדא	CARTOGAMO	GINGACCION	AAAACCTICTT	TTGCGMCC:	
	CTACTACTTO	AACTGCTGCT	CCACCTTGAC	GACCITCK:GAT	משכטלטטטטוני	COCTROCOLAT	GICACCITIC	CAGCTGCCCA	TTTTGCACAA	AACCCTGG()	
16301	GCACCACCGT	AGICTITACO	CCCONTRACC	CCTCCACCCG	CACITACAAG	CACGINITAIN	ATCAGGTGTA	CYCCGACGAG	GACCTOCTTO	ACCAGGCCAA	
	COTOOTOOCA	TCAGANATGC	AGACCACTOS	CGAGGTTGGC	CICCATGITC	CCCCACATAC	TACTCCACAT	OCCOCTOCTC	CTOGACGAAC	TCGTCCGGTT	
16401	COARCOCCTC	GOCCACTTO	CCTACGGAAA	REGGEATANG	CACATRICTAG	いっていっていること	GGACGAGKGC	ANCCCANCAC	CTAGCCTAAA	GCCCGTAACA	
	OCTOGOGGAO	CCCCTCAAC	COATCCCTTT	COCCGIATIC	CTGTACGACC	GCANCOGCGA	ccracreces	TICOCTICTO	GATCOGATTT	COCOCATTGT	
	P. 21									Kynt	
16501	CTUCAGCAGG	NOCTOCCOC	OCT TOTACO	TCCGAAGAAA	Accedence	ANAGREGAG	TCTOSTGACT	TOOCACCCAC	COTOCAGCTO	ATCCATACCCA	
16601	AGCCCCAGCG	ACTOGAAGAT	GICTIGAMA	AAATGACCCT	GGAACCTRAX	CTEXANGECEG	AGGTCCCCOT	GCGCCCAATC	ANGCAGGTGG	COCCGOCACT	
	Tracactrac	TOACCTICTA	CAGAACCTTT	TTTACTGGCA	CCTTRIGACTC	CACCTCOOGC	TCCAGCGCA	CGCCGGTTAG	TYCOTCCACC	GCGGCCCTGA	
16701	OCCUTOCAG	ACCORDOACO	TTCAGATACC	CACTACCAGT	AGCACCACTA	TYCCACCGC	CACAGAGGGC	ATCGAGACAC	AMCGICCCC	GGTTGCCTCA	
	CCCCCACOTC	TOCACCTOC	AAGTETATOG	GTGATORITCA	TCGTGGTCAT	AACGGTGGCG	Greatcacco	TACCTCTOTO	TTTOCAGGGG	CCAACGGAGT	
16801	accordeced	ATOCCOCOOF	CCACCCCCTC	OCTUGOGGCG	CGTCCAAGAC	CTCTACGRAD	GITTCAAACTR	ACCCGTGGAT	GTTTCGCGTT	TCAGCCCCCC	
	COCCACCOCC	TACOOCOCCA	COTCCGCCAG	CGACGCCCCC	CCAGGITICIG	GAGATACCITC	CACGTTINACC	TOCCCACCTA	CANAGEGERA	ACTCCCCCCC	
16901	000000000	CCGFTCGAGG	AMGTACGGCG	CCGCCAGCGC	GCTACTGCCC	CANTATRICCE	TACATCCTTC	CATTGCGCCT	ACCCCCCCCCT	ATCGTGGCT	
	CCCCCGGGCGC	OCCARGCTCC	TTCATGCCGC	GCCCGTCGCG	CCATCACCOC	CTTATACGES	NTCTAGGAAG	GTAACGCCGGA	TOCOCCCOA	TAGCACCGAT	
17001	CACCTACCGC	CCCAGAAGAC	GAGCAACTAC	CCGACGCCGA	ACCACCACTG	CAACTCGCCG	CCCASCATICGC		CCGTCCTTAGC	CCCGATTICC	
	OTOGATOSCO	OCCUPATION	CTCCTTCATO	GGCTGCGGCT	TRGTGCTGAC	CTTOSCOCC	DOCCOCKAGCG	GCAGCGGTCG	GOCACGACCO	GCCTANAGC	
17101	CTCCCCAGGG	TOCCTCCCGA	AGGAGGCAGG	ACCUINGING	TRECONCARC	CHOCTACCAC	CCCARRIATEG	TTTAAAAGCC	concrincia	GTYCTTGCAG	
	CACGCGTCCC	ACCCAGCGCT	recreecence	TEGGACCACG	ACCASITICACO	CCCCATGGTG	GOCTOCTAGE	AAATTTTCGG	CCAGAAACAC	CAAGAALGTY	
										luis:	
17201	ATATOGCCCT	CACCTOCCOC	CHCCGTTTCC	COCTROCTOCO	ATTICCGAGGA	ATTICIONORIA AGANTIGCACO GTAGGAGGIGO CATGGCCGGC	GTACAAGGGG	CATOGCCOGC		このこのことは、このこのこのこと	
	TATACCOCCA	GTOGACGGCG	GAGGCANAGG	OCCACOCCCC	TAMASICTICCT	TCTTACGTGG	CATCCTCCCC	GTACCOGCCG	GTGCCGGACT	OCCCCCCCTA	
	翻				Sphil						
17301	OCONCOTOCO	CACCACCOGC	3333333333	GTCGCACCGT	CCN: ATM SCINCIG	CCCCTATCCT	מככנכונכונו			CATTGREGGC	
	COCAGCACGC	GTOGTOGCCG	ככפככפניפכם	CAGCGTGGCA	OCCITACOCYC	CGCCATAGGA	CCCCCCA	TAAGGTGACT	AGCGGGGGCG	CTAACCCCCI	
17401	GTOCCCCCAA	TTOCATCOOF	GGCCTTGCA 0	GCGCAGAGAC	ACTUATTANA	AACAAGTTAC	ATCTCCAAAA	ATCANATA	MAGTETOGA	CTCTCACGET	
	CACOCCCTT	AACCTAGGCA	CCGGMCGTC	COCCIECTOR	TCACTAATT	THEFTENCO	TACACCITIT	TAGITITATI	TTTCAGACCT	GAGAGTECETIA	
										Econ TV	
17501	COCTROOTCC	TOTANCTATE	TTCTAGAATG	GANGACATCA	ACTITICOGIC					TYSCAACATA	
	GCGNACCAGG	ACATTGATAA	AACATICTTAC	CTTCTGTAGT	TCAAACGCAG	AGACCTAROLIC	CCTGTGCCCA	GCGCCCCCCAA	GTACCCTTTG	ACCOFTICITATI	

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		GUTGUCGCUSA MANTICTUSAS TCTACGACTC AGTGCAAAAT TCACGTTTTA CGTCCCCCCCG GCAGGCCCCG	* •			- · ·			GCAGLANGO" TAGCGGCCN: ATCGCCCCN: TOTCACAGN: ANTICOTIC: ATTTCOTIC: CACTLAGCN:
TOGGCAGGC TGCCAGGCC ACGGTCCGG CTGGCAAAG GACCGTTTC GCGTCCATG	E UDUDES	-			COCONCINATOS COCONCICOS CONTECTION GITACOSACT CICCOCOTTT GICIOGEOGRAMA	GCATTGCGAC CGCCGCGACG GCGGCGGACG AGCGCCGACG TCGCGGCTGC CCAACATTGGC GGTTCTACCG		Ped OCCUNTEGATO CCCTAGEAAC TAGETAACOT ATCATTGCA ATCATTGCA	COCCOOTAG COCCOOTAG GCCCOOTATG GCCCOOTATG GCCCOOTAG GCCCCCTA AGTOCTCTTA TCACCAGCTTT
TCGGGCCAGG AGCCCGGTCC CGGTGGCGCG GCCACCGCGG GCCACTCACC CGCCAGGTGCCC CGCCAGGTGG TACTCTGCCA ATGAGACCGT		ACCCTCGGA TOCOGAGCCT TACCCACGAC ATGCOTGCTG CTACCTCTGG GATCGACAC CTGCCTACAA	GTACCTGAGE CATGGACTEG GTGACCACAG CACTGATANCEG CACTATTCGC CACCTTGACT CCCCTTGACT CCCTTGACT CCTTGACT CCCTTGACT CCCTTCACT CCTTCACTTCA	CCCCAGCTGG GAGCCCGACC ACCCAGAGT TGTGCTGGAC ACACCAGATTG CCCAAGATTG CCCAAGATTG	TOCKATTING ACCITANACG CCCTTYCACG CCCANACTIC ATOCCTTCCA TACCCANATCC CCCCANATCG GROGITTANG	CCGCGCCANC GCCCCANCT GACGCCANGT CGTACTTICA GCATGAAACT TYGCGAATGG		AGEGGACTE CCOTGAGGAT GCCTCCTA GTCCTCGACC CACGACCTCT CTACTGCTCT GATGACGAGA	ATTOTTCAAA ACTGCGTACT TGACGCATGA GGGGCCCTA CCCCGGGAAT TGAAATAAA ACTTTAATTT
AGGACTACT CCCTGCTACT CCCATAGGT ATTAATCATG TACTTGTAAA AAGAACATTT TAAAGGGGTA		CAACGAAGAC GTTCCTTCTG ATAGGTGTCG TATCCACAGC CAGCTGGGAG GTCGACCCTC GCAACAAAAT CGTTGTTTTA AACATGTCAC	GAAGTAGACT CTTCATCTGC AAGGTCAAAC TTCCAGTTTG AGGAAAACTTT GGAAAAGCTAG CCTTTTGATG AAGATGTAGA	TEGETECACT TEGETECACT TEGETECACT TEGETECACT TEGETECACT TEGETECACT TECTGATEG ANACTICATICA TETTCATTICA TETTCATTICA TATAGATACC ATATICTTICG	GCCGATTITT GCCGATATATA GCCGATGAAAC CAATGAAAC GTTACTITGG GTTACTITGG CCTTTACGTT CCACACACTC	ACTCACGTAT TGAGTGCATA CATTTCAACC GTAAAGTTGG AN :TTACGGT TACAATGCGA TTTTTCTCAA AAAAGAGT ATATAAAAGAAT TATAAAAGAAT	TTCCCCAGC AACCCGTCCG TCAACCTCAA ACTTCGAGTT TCATATGCAA AGTATACGTT CTACTGAGGC GATGACTCCG CATGCCCACT GTACGGTGA	CCCTTATTCT CCGAATAAGA ATACCACCTTA ACCCACAAA TTCGACGACTTT ACCCCCACCAC TCGACGACCCC TCGACGACCCC	CCATATTANT CCATATTANT CTCAGTYSTA GAGTCACCAT TOAAANTCG* AATCGTGAT TTACCACTAY GTAACTCACG; CATTGATTG;

Figure 15L

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10101		TAN A CANA	TAPATCICE AB	Characteriant	TACATTIGETT	TTACKER 1	TTTTATTCATT (CTANTGRATT	ACAACAGCAC	GOSTAATATG
10561	TO THE PARTY OF TH		CATACOSCIT			AATCCCTGTT /	NANATANGGA	GATTACATAA	torrotecto	CCCATTATAC
	10 10 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1		Arberta Armen	A A TY S THE TENTS		אנאנאנאנאאני ז	ACAGAGCTTT	CATACCAGET	PPICCTICAT	TECATION
19401	CCACABERGE	-	TAGGGTCAAG	TTACKSACAAC	•			GTATGOTCGA	NANCGAACTA	ACCTARCCAL
. 6 2 6	CONCREDE		Particle hatch	A STATISTICAL AND A STATISTICA		CACATEITTA C	GANTTATTGA	ANATCATOGA	ACTORAGATO	AACTICCAAA
10661	TATACARCAG	CATCA	TACACCTTAG	TCCGACAACT	_	_	CTTAATAACT	TETAGTACCT	TOACTICTAC	TTCAA: x:1717
19601	TACTOCITY	_	GICTGATTAA	TACAGAGACT	•	_	אאכאמטשנאפ	CAMATGGAT	GOCANANGA	TCCTACAGAA
	AATGACOAAA		CACACTAATT	ATCTCTCTGA	GANTOSTICC !	ATTERCATE !	rigiccaurc	CITITACCTA	CCCTTTTCT	ACGANGICIT
19701	TITTCAGATA		AAGAGTTCGGA	AATAATTTE	CCATCACAAAT	_	CCCAACCTATE	CCACACAATTT	CCTOTACTCC	AACATAGCC
	AAAAGTCTAT		TTCTCACCT	TTATTANAC	GGTACCTTTA O	GTTAGATTTA	COGFICCACA	CCTCTTTAMA	GCACATGAGO	TGIANCA
19801	TOTATITICC	CCACAAGCTA	AAGTACAGTC	CTTCCAACGT	AAAAATTTCT	_	ACACCTACCIA	CTACATGAAC	ANGCGAGTOG	TOGCITCCC
	ACATANACOO	-	TTCATGICAG	CANCETTOCA	TTTTTAMGA	CTATIOGRAFE	rorcoatoct	CATCTACTTO	TICOCICACE	ALT GAGGG.
19901	CCTACTOGAC	TOCTACATTA	ACCTTGGARG	ACCCTOSTICC	CTTGACTATA '	PATACAACGT (CAACCCATTT	AACCACCACC	CCAATGCTOG	cerdeactw.
	COATCACCTO		TOGANCCTCG	TGCGACCAGG	GAACTGATAT	ACCTICITIGGA (STIGGSTANA	ricorcorda	CONTRACGACC	GGACGCGATT
20001	CRETEABILE	-	TOGICACTAT	GIGCCCTICC	ACATACANAT	GCCTCAGAAG '	TTCTTTGCCA	TTAMANCCT	CCTTCTCTG	CCGGCTCAT
	COCHETTACA		ACCAGCGATA	CACCICIONAGG	TGTANGTCCA	COCAGTCTTC	MGNACGGT	AATTTTTGGA	GGAAGAGGAC	GGCCCGAGTA
					Pstl					
•0.00	State	Control & Control	ACCARCERITE	TENACATICAL	TCTTSCAGAGG	TCCCTAGGIA	ATCACCTAAG	COTTCACCCA	CCCACCATTA	AGTTTGATA:
70102	TOTAL STATE		TCCTTCCTAC	AATTGTACCA	ACACTITETE	AGGATCCTT '	TACTOGRAFIC	CCAACTGCCF	COCTCCTAAT	TCANACTA'R!
			distribution of the later of th	COSTOCACAAC	ACCCCCCA	CCCTTGAGGC	CATGCTTAGA	AACGACACCA	ACCACCAGTC	CTTTANCO!
70707	CATTIGCCIT		AGAAGGGGTA	CCCCCTGTTG	TCX XCX CACCT		GTACGAATCT	riccrotoor	TGCTGTTCAG	DAMITICE :
	Olomoro de		Carrie Transcript	ATACCCCCA	ACCTACCAA	COTGCCCATA	TCCATCCCCT	CCCGCAACTO	OCCOCCITIVE	_
70201	TATCICICO		COACATOOGA	TATOGOCGOT	TESCATEGIT	CCACCACTAT	ACCTACCCCA	GOCCOTTGAC	CCCCCCAAAG	_
	A I RUMONOMON		ANGRANACTO	CATCACTGGG	CTCCGGCTAC	GACCCTTAIT	ACACCTACTC	TOCCICTATA	CCCTACCTAG	_
70407	GOANGIGGG		TICCTITIOGO	GTAGTGACCC	GAGCCCGATTS	CTCGGAATAA	TCTCCATCAG	ACCGAGATAT	GGGATGGATC	
10201	der betreet being		AGANGGTGGC	CATTACCITT	GACTCTTCTG	PCARCTRACE	TECCHATCIAC	CCCCTCCTTA	CCCCCAACGA	-
1000	AATGGAGTTG		•	GTAATGGAAA	CTGAGAAGAC	NOTICANCEDO	ACCGITTACTG	GCGCACCAAAT	COCCUTICAT	CANCELLIAN
20501	AACCISCICAG		GGGTTACAAC	GTTGCCCAGT	CTAACATGAC	CNANGACTEG	TICCTIFICAC	ANATOCTAGE	TACTATANC	TABLETACE
	Treacanate		CCCAATGITTG	CAACGCCTCA	CATTIGTACTG	GITTCICACC	MAGGACCATO	TITALICATIC	AI IGAINA IA	•
20701	ACABACTICTA		ACCTACAAGG	ACCCCATGTA	creentern	AGANACITICO	AGCCCATGAG	CCGTCAGGTG	GTGGATGATA	
	TCCCGAAGAT	_	TCGATOTICC	TYCCGTACAT	CACCOAAGAAA	TCTTTCAACG	TCGGGTACTC	GCCACICCAC	CACCIACIA	
10801	GOACTROCAA	A CAGOTOGOCA	TCCTACACCA	ACACAACAAC	TCTOSATTUS	TTGGCTACCT	יוטכניכככאככ	ATOCOCOANO	GACAGGCCTA	
1000	CCTGATOGIT	-	-	rotement	ACACCITAAAC	AACCCATOCA	ACCICACIONO	TACCACACTIC	CIGICLOGA	
							***************************************			Try. Marth After
20901	TECCCTATE	C COCTTATAGG	CAAGACCTACA	CETTGACAGCA	TTACCCAGAA	AAAGITHCTT	TYCCEATCOCA	CCCT11GGCG	GTAGGGTANG	
i i	AAGGCGATAG GCGAA	G OCCUANTATOC	GENCHORICGE	CAACTCATCGT	AATGGGTCTT	TTTCMMGAN	ACTACTACACT			

Figure ISM

_:	GNOSTICIATE CCATGRANGA CTCCACCTAG OGIACCTCCT	CACGOCCTT : GTGCGGGA/ 1: Ingili	\$ F				CACCATGAA!		_		ACTORISOTOR		CACGAGTCC		· ircrcaa	GGCCCCACCG GTTCTTCACG CCGCGCTCGC CAACAAGTCC
Ramel	GNOCHEGATE	TOTACCTOCG AC/ "YGACGC	AAAACCATTO	TAGTCAATAC ATCAGTTATG	ANDACTOOTC	Pre AGG TOO	ATCACAACCC TAGTGTTGGG	-			CCACTTCAAA			GTAGTTTTCC AAGAACATGC	TTCTTGTACG	GGCCCCACCA
	CATCACTETT	ATCGAAACCG TAGCTTTGGC	GCAGGAACTG	GCCTGCGCCA CGGACGCGGT	CCTTROCCTT	AACGCTGGAA	ACTCCCATGG TGAGGGTACC	AACAGCTCTA	AAAATAATGT		CGCCGTCGAG		-	ACCOTAGNOS TOCCATOACO	COGNACI: TC	ACCACATTIC TOCICITAAAG
	ACGCGCTAGA TGCGCGATCT	GRICACCOUNG	CCTCCAGTGA	ACACAAGCTC TGTGTTCGAG	CTCTTTGAGC	ACCOCTOTAT	CTOCACCCCAA		ANAACATGTA		ACAACCATCC TGTTGGTAGO		_	TTGCACTCGC AACGTGAGCG		4 95 5
	AACTKSCRICK'C T'KGACK'SCRIG	ACCAPTCRCA:A TUGICACCA	GCCCCATCC	TRITTACTCC	MCATGCTAC	TCTTCCCCCG AGANGEDGGC	CCTTTGCCAA	CACCCTGCGT	TOTCACTTGA ACAGTGAACT	THICCGRETO AACGCCAGAC	AAACTCAGGC TTTGAGTCCG	AAGTCGCAGT TTCAGCGTCA	CCCACATACAG	ACCOMMETE AAACCAAACTC	TTTCGGTGGA	
:	TCTCTACOTC	GAGGGAGAGAGG	AACAACAGT	TTTCCAGGCT AAAGCTCCCGA	CGCACTENAN	CGCCATTGCT	TTTCTCCACG AAAGAGGTGC	AGGTACAGCC TCCATGTCGG	CACTICITIT	ACCCCCACCC TXXXXXCTGGG	ACTACCACTT ACTACCTIGAA FMRV	CCATATICITIC	CHACCAGGA	COCCICATION	CTAGACGAAT	GREGITICINES CARCACETTIS CAGCACUTICE GIEGITISAAC
	COSTITIONS		AACCAACATC	TTACAAGCGC ACTGTTCGCG	GCCTFSGAACC	TCCGCCGTAG ACGCGGCATC	CTYSCTISCATIG	AACAGTCCCC TTOTCAGGG	TTAGGAGCGC AATCCTCGCG	GTGATTATTT	TOGTGTTTAG ACCACAAATC	CCAGCCCCC	GTGGTGCACG CACCACGTGC		ATTITICGGAA	GACAGGCGC GACGACGACG CAGGACCTTG
	ACAGACCTGG		ATMACAAGC		CTACCGGAAA	CHCAGTGAGG		CTCCATGCTC GAGGTACGAG	AGTOCCCACA TCACGCCSTCT	ACACTCTCGG TOTOAGAGCC	OTTGCCATAC CAACGCTATG	OCCITITACCA CGCANATCOT	TCAGCGCC08 AGTCGCGGCC	TAGCTGCCTT ATCGACGGAA	MIGGACTITICA TCCCCAACGT	GACAGGCGC
	COCCACTC		ACCCCACAAC		CCCATGTGAC	GATTONOTAC GAGICACTCC	TCOOCCOCCT	CCCATOOOTT	CCCCAGCCAC	TITTATTIOT AMATAMACA	OCAGGGACAC COTCCCTGTG	CATCACCAAC GTAGTGGTTG	TOGAACACTA	TCAACTITOG AGTIGAAACC	CAATCCTATG	GCCGRAAAC TGATTGGCCG CGCCCTTTTG ACTAACCGGC
	TTATOTCCAT		TCOCCCOCA		ONDACTODOS CTCTGACCCC	AGGITTACCA		CTTATTACCO	CCCCTACTT	AGCANATOC	TOCOCCACTO ACOCCOTCAC	GOCTGCOCAC	GTTGCAGCAC	OCCANCOCKO COCTTOCCTC	COCTCTOOOC	GCCGGAAAAC
	21001	21101	21201	21301	21401	21501	21601	21701	21801	21901	22001	22101	22201	22301	22401	22501

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ישינ	da de la constante de la const	THE PROPERTY.	لطالاتطاطة	COCOLAMACIC	T. P. L.	CHITCHEATEC !	ATTICANTO	CONOCICCIT	ATTTATCATA	ATGCTTCCGT
10077	ALL HOUSE	TOC PROPERTY.						GCACGAGGAA	TAMATAGTAT	TACGNAGGITA
	TAGAALCIAGA	ACGAICIONC								15.1
•						י בייני	accountable.	ATCCTTCTAG	GICACCICIN	CNANCGACTO
10/22	GTAGACACTT AAGCT	MACTICALLI	TCGRITTING						2424004045	Contento (Colic.) At
	CATCTGTGAA	TYCOAGCOGA	AGCTAGACIC	CCCTCCCCAC	ניותאאייהדוה	כטייין ביינולכי	ACCEGAGEAC	TACGAACATC	CAGILCCACAC	GITTI CHA
	P.S.II	Pstl								
22801	TACOC		OCCUCATICAT	CCTCACAAAG	GPCTTRITTEC	TEMPTOTATION	CAGCINGDAAC	CCGCGGTGCT	CCTCGTTCAG	CCAGOTCT'1.1
	GTCCATGCCC		CGACGTAGTA	GCAGTGTTTC	CAGAACAACG	ACCACTTORA	GTCCACGFFG	GOCCCCACGA	GCACCAAGTC	GTTCCAGAAC
22901	CATACGCCC		CACTICATICA	CCCAGTAGTT	TOAAGTTOGC	CTITAGATED	TTATCCACGT	COTACTTOTO	CATCAGCGCG	CYXCYCAGCC'F
	GTATGCCOGC		GTGAACCAGT	CCÓTCATCAA	ACTICAAGCG	GNAATCTAGC	AATAGGTGCA	CCATGAACAG	GTACTCGCGC	acaconca 1
			Fvid							
23001	CCATGCCCTT	CTCCCACGCA	GACACGATCG	GCACACTCAG	CGGGTTCATY:	ACCOUNTER	CACTITICCGC	Treastroope	remecter	CCTCTTGCC1"
	OUTACGODAA		CTGTGCTAGG	COTOTOAGTC	CCCAAGTAG	TOCCATTANA	GTGAAAGGCG	AAGCGACCCG	AGAAGCAGAA	CCAGNACGCA
21101	CCCCATACCA		CONTROLLE	ATTICACTOC	CCCACTGACC	GCTTACCTCC	TTTRECEATER	TTCATTAGCA	ccoorecon	GCTGAAACCC
	OCCUTATOGT		CCAGCAGAAG	TARGECTOCC	GCGTGACACG	CONNTRACAGO	MACGCTACG	AACTAATCOT	GCCCACCCAA	CCACTTTCO
23201	ACCATTIGIA		TETETETET	TCCTCXCTGT	CCACGATTAC	CTCTCTCATGAT	GGCGGGCGCT	COCOCITICOC	ACANGGGCGC	TICTUTION
	TCCTAAACAT		AAGAGAAAGA	AGGACTICACA	CHICCTANTO	GINGACICACTA	CCGCCCGCGA	OCCCOANCCC	TCTTCCCGCG	AMGAAANNGN
23301	TETTRACECOC		٦٠٥٥٥٥٥٥	ACCTOUNTED	CCACACACTO	CONTRACTOR	GCACCAGGG	GICTIOTOAF	GAGICTICCE	CCTCCTCGGA
	ACAACCCCCC		AGGCGGCGCC	TCCAGCTACC	B BCGCCCGAC	CCACACGCGC	corocreces	CAGAACACTA	CTCAGAAGGA	GCAGGGAGCCT
21401	CTTTATACAC	CACCICATOC	0011111100	0000000000	מטעטטטטטטט	CCCACCOCGA	CHARGACGAC	ACCIECTECA	toerrecese	ACCTCGCGG.
	CACCTATGCG		CGANANANCC	CCCCCCCCCCCC	CCFCCCCCCCC	CACTGCCCCT	accer:10c10	TCCAGGAGGT	ACCAACCCCC	TOCAGCGC
21501	G.ACTORITY		GOTGOTTICO	COCYCCTCCT	CTTCCCGACT	CACCATTACC	TYCTICTATA	GOCAGANANA	GATCATOGAG	TCAGTCGAL:A
	CCTOCCCAG		CCACCAAAGC	GCGACCAGGA	GAAGGCTGA	CCCCTAAAGG	ANGAGGATAT	CCONCINITY	CTAGTACCTC	AGTCAGCTCF
23601	ACAAGGACAG		CCCTCTGAGT	TOCCORCAC	CULTICCACC	GATGCCGCCA	ACGCCCCTAC	CACCITICCCC	GICCOROGEAC	CCCCGCTTGA
	TCTICCTOR	CLADO	GGGAGACTCA	ACCCCTCCTG	GCIXCACASTAG	CTACGGCGGT	TCCCCCATG	OTOGNAGOGO	CAGCTCCGTG	COCCCCAACT
21701	Actionation		AGCAGGACCC	AGGEFFFETA	NGCGNAGACG	ACGAGGACCO	CTCAGTACCA	ACAGAGGATA	AAAAGCAAGA	CCAGGACAN'
1015	CCTCCTCCTT		regreerood	TCCANNACAT	TOCOTION	TOCTCCT/GGC	GAGTCATOOT	TOTOTOCTAT	THICGING	CONCERGIANS
1005	Agreement of the Control of the Cont	ACCRETABLE	ACTURATION	GGGGACGAAA	GOCATOGOGA	CTACCTAGAT	GTOGGAGACO	ACGRECATIT	GAAGCATCTG	CAGCGCCAGT
70067	Contraction of the contraction o		TEAGCCCOCC	CCCCTGCTTT	CCCTACCGCT	GATCGATCTA	CACCCTCTOC	TECACGACAA	CTTCGTAGAC	GTCGCGGTCA
11001	COUCERTER		THEMONGO	GCACCCATGT	מכבכבעבעב	ATAGCGGATT	TCAGCCTTGC	CTACGAACGC	CACCTATTO	CACCOCCCT
43301	CECCENTARTA	GACOC	AACGITICITCO	COTCOCTACA	CHANGEARCORD	TATCCCCTAC	AGTICGGAACO	GATOCITICO	GTGCATAAGA	OTCOCOCOCA
******			ACCAPACATO	CCACCCAAC	CCCCCCCCACA	ACTICIACCC	CGTATTICEC	GTCCCAGAGG	TCC TTCCCAC	CTATICACATIC
T0057	ACCCCC		Transfer AC	0010000110	COCCCCCACT	TCAACATGEG	GCATAAACGG	CACGGICTCC	ACGAACGOTO	GATACHCITA'S
		3								Ecosto
		TADA STATES	ACTOR OF A TOP	Trachettecta	VECCENTED	AGCGGACAAG	CAGCTCCCCT	TREGGEAGG	CGCTGTCATA	:כנדעואדאדה:
10162		AAAAAGGTTT TGACGTTCTA	TOGOGATAGG		TGGCGTCGGC	recentation	GTCGACCGGA	ACCACCATACC	GCGACAGTAT	GGACTATAGY.

Figure 150

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24201	CCTCGCTCAA CGAAGT	CGAAGTCCCA	AAAATCTTTG TTTTAGAAAC	ACCCAGAACC	ACGCCIACTAG TGCGCTGCTC	AARTONIATO	CANACCICITE	CCANCAGGAA	AACAGCGAAA	atgaaa;tca tact'ftcagt
		\$	Xhol							
24301	CICIDOAGIO	TTCOTOCAAC	PROGRAC TOGACCITCA	CAACGCCCCCC	CTAGECGENE	TANAACTECAG	CATCGAGGTC	ACCCACTITO	CCTACCCCCC	ACTTAACCTA
	GAGACCTCAC	AACCACCTTO	AGCTCCCACT	GTTCCCCCC	GATCGGCATG	ATTENTION	GTACACTCCAG	TOXICHOMAN	GCATGGGCCG	TRANTIGGAT
24401	CCCCCCAAOO	PCATGAGCAC	AGTCATGAGT	GARCTRATEG	THEORYCOMOC	שנאפרנינדה	GAGAGGGATG	CAAATTIGCA	NONACARACA	GAGGAGGG
	OGOGGGTICC	AGTACTOGIG	TCAGTACTCA	CTCGACTARC	ACCATCACTACG	وروالان المعادد المداد	CICACCTAC	GTTTAMACGT	101101101	רוויני ווינינואי
24501	TACCCCCAGT	TOCCGACGAG	CACCTAGGG	GCTGACTTCA	AACGCGCGAG	CCTGCCGACT	TRIGARIGACIO	ACCCAAACTA	ATCATCCCC	CAGTN:CTCCT
	ATGGGCGTCA	Accectocic	GICGATCGCG	CCACCGAAGT	TYNCHOOCIC	CCACCICITICA	ACCTCCTCCC	TOCCTITICAT	TACTACCOCC	GTCACGAGEA
		PH-IS	## ***********************************							•
24601	TACCGTGGAG	CTTCAG	TGCAGCGGTT	CTTTGCTMAC	CCGGAGATGC	AGCGCAAAGCT	AGACGAAAGA		CCTITICGACA	GOCCTACGTA
	ATOCCACCTC		ACOTOCCCAA	GNACCACTG	GGCCTCTACG	TCGCGTTCGA	Terecrings	AACOTOATGE	COMMECTOR	CCCGATCCAT
		Both								
24701	CACCAGACCT	8	CAACOTGGAG	CTCTGCAACC	TRESTUTECTA	CCTTCCAATT	TTGCACGAAA	ACCOCCTTGO	GCANAACGTG	CTICATTCCA
	GCOCHCCOOA		GTTCCACCTC	CACACCTTOG	ACCNGAGGAT	GGAACCTTAA	AACGTOCTTT	TOCCOGNACC	COTTITICCAC	GAAGTAAGGT
		Ascl	3							
24801	CGCTCAAGGG	g	COCGACTACO	TCCGCGACTG	CCTTTACTTA	TERCTANGET			GOCOTITIOGIC	AGCAGTUSCTT
	CCCAGTICCC		GCGCTGATGC	AGGCGCTGAC	GCMATGAAT	AAAGATACGA	Trytoxiaccet	Croccootac	CCCCCANACCO	TCGTCACGAA
			Psil					!		
24901	GCAGGAGTGC	AACCTCAAGG	ACCTUCANCIA	ACTURCITABAG	CANANCTIGA	ACTACCTATG	CACCACCTTC	AACGAGCGCT	CCCIIOCCCOC	GCACCTGB(7:
	CCTCCTCACO		TCGACOTCTT	TGACGATITIC	GTTTTCAACT	TECTEGATAC	CTGCCGGAAG	TRICTICGEGA	GGCACCCGCG	CCTCGACCGC:
25001	GACATCATTT	TCCCCCAACG	CCTGCTTAM	ACCCITICAAC	ARKINCE	AGACTTCACC	ACTCAAAGCA	TCTTCCAGAA	CTTTAGGAAC	THE VICCIA.
	CTOTAGTABA		GGACCAATTT	TRACARCETTE	TECENGACIO	TCTGAAGTGG	TCAGITTCGT	ACAACGICIT	GANATCCTTG	AAATARGATK'
25101	AGCOCTCAGG		GCCACCTGCT	GTGCACTTCC	TAGCGACTIT	GIGCCCATTA	ACTACCGCGA	ATGCCCTCCG	CCGCTTTGGG	GCCACTOCTA
	TCGCGAGTCC	TTAGAACGGG	COCTOCACGA	CACGTGAAGG	ATCCCTGAAA	CACGGGTANT	TCATGGCGCT	TACGGGAGGC	GCCGAMACCC	COCHEMETERY
	the part									
25201	CCTTCTGCNO	CTAGCCAACT	ACCITIOCCTA				TGACOSTCTA	CHAGAGRETC	ACTORCOCTO	CAACCTATA
	GGAAGACGTC	GATCGGTTGA	TOGAACCCAT	GGTGAGACTO	TATTACCTTC	TOCACTOCACT	ACTOCCAGAT	GACCTCACAG	TGACAGCGAC	

25301	ACCCCCCCACC	acrecetos?	TTOCAATTCO	CACCTCCTTA	ACCIMAGICA	AATTATCGGT	ACCTITICAGC	TOCAGOGICC	CTCGCCTGAC	GANAGICCE
	TOCOCOUN	CGAGGGACCA	AACCTTAAGC	GTCGACGAAT	TCCTTTCAGT	TTAATAGCCA	TOGRANCITCO	ACCITCCCAGG	CACCCCACIG	CHILINGS
25401	CONCECTO	_	ACTCCGGGGC	TOTOGACGTC	COCTITACCTT	COCANATITES	TACCTGAGGA	CTACCACGCC	CACGAGATTA	COTTCTACGA
	GCCCHOCCC		TGAGGCCCCG	ACACCTGCAG	CCGNATGGAA	CHICATTIANAC	ATOCACTCCT	GARGORGEGG	GTGCTCTAAT	רכאמישומ
25501	MANCCANTCC	COCCCOCCTA	ATOCONACT	TACCARCTUR	CHIATTACCC	ACCOUNTANT	TCTTICATICAA	THECHAGECA	TCAACAAAGC	CCCCCANON
	TCTCGCTTAGG		TACCCCTCGA	ATCCCCCCALT	CACTAATAGG	TOCCOMMETA	ACAACCOGTT	AACGTICGGT	ACTIGITION	ואירואיזורזי
25601	TTTCTCTAC	GANAGORACG	GOOGSTITTAC	THYCACTCCC	ACTROCOCICA	CCACCTCAAC	CCAATCCCCC	COCCOCCOCA	CCCCTATCAG	CARCAGRECA
i i	AAAGACGATG		CCCCCAAATO	AMPLITATION	TCAUCCCUCT	CCTCGACTTG	CONTINGENCE	GCCCCCCCC	CCCCATAGTC	Greener



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25701	GGCCCTTGC	GOCCCTTGC TRCCCAGGAT	GCACCCANA	AAGAAGCTUC	N.X.Trayreacc	נשיגעינעניפ	CACGAGGAGG	AATACTGGGA	CARTCAGGCA	CAGGACACTET
•	CCCOGGNACG	CCCOGGACG AAGGGTCCTA	CCGRARATT	TTCTTCGACG	TYTHATACAG	CINTRAGENCE	CTGCTCCTCC	TTATGACCCT	GTCACTCCGT	CTCCTCCN .
						Harifff				
25801	TOCACGAGGA	CCACCACCAC	ATCATCANG	ACTOROGAGAG	נוניבאינאיניאינ	מאחנידדכנה	AGGTCGAAGA	GETGTCAGAC	GAMACACCOT	CACCE-Tecs .
	ACCTGCTCCT	cerecteero	TACTACCTTC	TGACCCTCTC	GGATIC TGCTK:	CITTCGAAGGC	TCCAGCTTCT	CCACAGICIO	CTTTGTGGCA	GTGGGAGCC A
25901	CGCATTCCCC	Traccoacec	CCCAGAAATC	GCCAACCGGF	TECHCEATEG	CTACAACCTC	CACTCCTCAG	GCCCCGCCGG	CACTGCCCGT	TORCORACCC
	CCCTAACCCC	AGCGGCCGCG	GOTHETTING	CCGTTGGCCA	ACKITCGTACC	CATCITISCAC	CCGAGGAGTC	cacacaacc	GTCACCCCCA	ACCOCTOCO
26001	AACCGTAGAT	GOGACACCAC	TOOMACCARG	OCCUGITANGE	CCANGCAGCC	CCCCCCCTTA	CCCCANGAGC	AACAACAGCG	CCAAGGCTAC	COCTCATEG!
	TTOOCATICEA	cccrcrccra	ACCTICKSTCC	CGGCCATTCA	GSTICGTCGG	CRACARCANT	CCCCTTCTCC	TIGHTOROGO	GGTTCCGATG	OCCIMOTACCO;
26101	OCCOCICACAA	GAACGCCATA	GTTCCTTGCT	TOCANOACTO	TOCOCOCAAC	ATCTCCTTCG	CCCGCCCACTT	TETTETETAC	CATCACORCO	TOXIC: THEFT
	COCCCOTOTT	CTTGCGGTAT	CNACCAACGA	ACGITICITGAC	ACCCCCGTTG	TAGAGGAAGC	GCCCCCCCAA	AGANGAGATO	GTAGTGCCGC	ACCHGAAGG"
26201	CCOTAACATC	CTGCATTACT	ACCONCATCT	CTACAGCCCA	TACTRICACES	CCCACACACCC	CAGCHACAGC	ACCOCCACA	CAGAAGCAAA	מאכנואככפנוא
	OCCATICTAG	GACCITANTGA	TOCCAGTAGA	GATGTCGGGT	ATGACGTOGC	CACCOTCCC	Greenste	recedence	grenteamr	CCGCTGGCCT
26301	TAGCAAGACT	CTGACAAAGC	CCAAGAAATC	CACAGCGCG	GCARCAGCAG	GADGACKACC	ocrocorcia	GCGCCCAACO	ACCCUTATE	GACCCCCCARG
	ATCOTICTOA	GACTOTITICS	GGTTCTTTAG	GIGICOCCOC	corcorcorc	CTCCTCCTCG	CGACGCAGAC	CGCGGGTTGC	TTOCOCATAG	CROGGCGCTC
26401	CTTAGAAACA	GCATTITICC	CACTETRITAT	CCTATATIFIC	AACAGAGCAG	GROCCANGAA	CANGACCTOA	AAATAAAAA	CAGGICTOR	CCATCCCTCA
	CAATCTITUT	CCTANAAAGG	GTGAGACATA	CCATATAAAG	TREPOTORIE	CCCGGTTCTT	GETCTCOACT	TITATIMIN	GTCCAGAGAC	OCTACCCART
26501	CCCOCAGCTO	CCTOTA	ANAGCGNAG	ATCAGCTTCG	GCGCACGCTG	האהאההככפס	AGGCTCTCTT	CACTAAATAC	TOCCACIOCICIOA	CTCTTAAGGA
	GOOCGICGAC	GCACATAGTO	Trincocrac	TAGTCCAAGC	CGCCTYCCGAC	CTTCTGCGCC	TCCGAGAGA	OTCATITATO	ACCCCCCACT	GAUDANTICC .
26601	CTAGTTTCGC		•	CCCMANCTA	COTCATCTCC	ANCONCCACA	CCCGGCGCCA	GCACCTGTTG	TCAGCGCCAT	TATITAGCANG
	GATCAAAGCG	COCCANACIAC	TITAAATICG	CCCTTTTCAT	CCACTAGAGG	TCCCCCCTCT	OCCCGCOOT	CCTCGACAAC	Acresector	ATACTICGTTC
26701	GANATICCCA	COCCCTACAT		CAGCCACAAA	TOCHACTICAL	COCTGGAGCT	CCCCAACACT	ACTCAACCCG	AATAAACTAC	ATGARCCCC
	CFFFAAGGGF	OCCOORTGEA	CACCTCAATG	Greedight	ACCCTGAACO	CCGACCTCGA	COCCUTICAGA	TCAGTTGGGC	TTATTTCATC	TACTCGCGCC
		CONT			Đ	ECOTI				
26801	CHOCCEACAT	CTATAGGGGCC	CAGITICACIT	TACOCOCCCA	CCCAAACCCA	CCCAMACCCA ATTETECTOR	AACAGGCGGC THGTCCGCCG	TATTACCACC	ACACCTCOTA TOTOGAGCAT	ATACCITIAN TATTCCAATI
26901	PCCCCGTAGT	TOCCCOCTG	CCCTOGTGTA	CCAGGAAAGT	CCCCCTCCCA	CCACTGTGGT	ACTITECTION	GACGCCCAGO	CCGAAGTTCÀ	GATTIACTIANT
	AGGGCATCA	ACCOCCCCAC	GGGACCACAT	GGTCCTTTCA	CACCCACCACT	GUTTOACACCA	TCAACCCTCT	CTGCGGGTCC	GOCTTCAAGT	CFACTGATTO
27001	TCAGGGGCGC	AGCITICOGG	COCCUTICGE	CACAGOGING	ומפורהכרנהה	GCAGGGTATA	ACTICACCTGA	CANTCAGAGG	GCGAGGTATT	CARCTCAACT
	Agreceded	TCGAACGCCC	GCCGAAAGCA	OPGRECEACG	CCARCAGISC	CGTCCCATAT	TGAGTGGACT	GITAGICTICC	CCCTCCATAA	Greadfra.
27101	ACGAGICOGI	GAGCTCCTCO	CHICCICACC	CAGGCCTTACC	GACATTTCAG	ATCGGCGGCG	CCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC	ANGTANGED	CCTCCTCAGG	CANTCCTAAC
•	Pati	20000012								
27201	TCTGCAGACC	restoriens	AGCCCCCCTC	TOGAGGCATT	GGAACTCTGC	AATTTATTGA	GGAGTTIGTG	CCATCOGICT	ACTITIAACCC	CITCICOGIA
	AGACGACTOG AGCAGG				CCTTGAGACG	TTAMTANCT	CCTCANACAC	CGTACCCAGA	TGAAATTGGG	GAAGAGCCT

Figure 1501

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GCAGAGCAAAC CGTK:TCC:1T:1	ATATECIAGIG TATACETEC''	CONTRACTOR IN	AGNANTIA:A ICTITAAT"	NICTOTICS TAGACACO	ACCITACCT:		CANTACAT	GCCTGCTK! ?		ACCACAGN/ TGGTGTCT1'} TTACAGTT1'F AATGTCAAAA	CACCHARAT TACANAGGA ATGITTICA ACANATIA TIGITTAN IT
MAGTOCAGAG	CCCCACCATC	TAGTTGAGCG	TATTAATAC ATTATTTATG	TACTITITAC	ACACCACCC	AGTTATTOAG	ACTCTACOOS TGAGATOCCC	AAGGCTCGCC TTCCCAGCGG	ACTOCOAACO	TATAAAATGC ATATTTTACG GAGTATAATG CTCATATTAC	AGTATMOTT TCATATTCAA CTATATTAAA GATATMATT CTCCTTCCAA GACGAACOTT TCTATCTCCAA
ACTOANTOTT TRACTTACAA	CHITGANTIO	CCCCCCTGC	CACGACTCAT CACGACTCAT	CCTTACCTOO	CATCAGAAAA	COCACAGACC	ANTICAROCA	•	CTAGGITTAC	GCACCACTCT COTGGTGAGA TGACACTACA ACTGTGATGT	ATGAGCAAAC TACTCGTTTG GTACCCTACT CATGGGATGA GCTTTACTCG CGAAATGAGC AACAATTGAC
CACCACTACO ACTODATOTT CTGCCGATGC TGACTTACAA	ACTITICATA TEANANCENT	GTTTACCCAG	TCCCATCTCT ACCCTAGAGA	CCANGGGAA	TCACCTACTC AGTCGÁTGAO	AGACTTTTTC TCTGAAAAAG	GTTTATGAAC	ATACTAACOC TATGATTGCG	GTACATAATC CATGTATTAG	CCTANTRACTCA CGATTACTCA CCCACCAGG CCGTCGGTCC	TACCATGTAC ATGGTACATG GCTTTTGGTCT CGAACCAGA ACCACTAACT TGGTAACT GATTCCCCTG GTAAGGGGAC
ההאהתהמכת הביוהאתהכים	CACTACAGECAE	TCATTCGGGA ACTAAGCCCT (Igill	AGATCTTTGT TCTAGANGA	CCCAAGCAAA	CTCTCCCAGC	ACCOTAMACC TGCCATTTGG	CTACTGTCATO	CTTTATTCTT	AGATGATTAG TCTACTAATC	CGCAGCTGAA GCCTCGACTT TATGCTATTT ATACGATAAA	ACACGCTGTA TACAGTGCTC ATGTCACGGG ACCTAATGTC TCGATTACMG TCCGATTACMG ACGGGTTATGG
ACCITICATION TO THE	CTT/NOCCICAL GAAAN TARACIG	GCCCCTACATC CCCCCCATCGG	CVANTATAGE	TCTTCACCCG AGANGTGGGC	ACGAGAGAAC TGCTCTCTTG	CTACCGCCTG	ANARGEGEAG TITTECEGEGTE	TIGIGATICT AACACTAAGA	TCGCCACCCA AGCGGTGGGT	ATGITACATT TACAATGTAA GTATGCTGTT CATAGGAGAA	TTTATGAMA NAMTACTIT CTATCCTAAT GATACGATTA TANGTIACAA ATTCAATCT CCAGTAAAGG
CCTAACTITG	CCCACAACTC	CCCTCTCGAA	CCTAACCCTG	AACGCCACCO TTGCCGTGGC	GAGTGAGTCT	TYSCACCACAC ACGROGIGIG	GTATTAGGCC CATAATCCGG	ATTCTCTGTC TAAGAGACAG	AACGCTGGGG TTGCGACCCC	CCAGCCTGTA GGTCGGACAT AAATTGGCAA TTTAACGGTT	BEILTON COT ATCTITICEA COT ATCANAGET THE TECTOCACTE INDE ACCACTEGA INDE CATANTITA INDE CANACECEE INT TANACECEE TAN ATTROCOCOC
TCAATTATAT		TTACCGCCCA AATGGCGGGT	TTGCAACTGT AACGTTGACA	CCATCCTGTA	AACCCAGACG TTGGGTCTGC	CACCOOCCGC	ACCCTTAGG TTGGGAATCC	OCPTGGGGTT ECAACCCCAA	CACCTTTTTA	TTTTAAGGAG AAAATTCCTC CACAAAAACA GTGTTTTTGT	CTTTTAT GAAAATA TGGCACT ACCGTGA AAGAAAA TTCTTTT GAATACC
ACTATCOOM		GOCUTECCOC TTACCGCCCA CCCCAGGCCG ANTOGCGCCCT	TCACTOTGAT	OCTCCTATCG CGAGGATAGC	CAACAGITIC	ACCIACIOCOT	GAGCTTAGAA	The CANGAATCGG	CATTIATIOT GEAAATAACA	AAAAGGTGGA TTTTCCACCT GCTTATTCGC CGAATAAGCG	AUTCAFAAAA TCAGTATTTT TGGAAAACAC ACCTTTTGTG TATTGAGGAA ATAACTCCTT ATTATAATTA
CCICCCOOCC			CCCTOTATIC	ATATACTOOD TATATGACCC	CTOTOATTTA	CCGGGAACOF	ACAGGAGGT	TCKGOTTPCT CTAGAATCGO	TOCACATTEG	GOTACCACCC CCATGGYGGG ATGAAAAGCT TACTTTTCGA	CCAGGGTAAA GOTCCCATTT CAAATTTAACAC GACTCAGCTF CTGCGTCGAAAAAATTTAGCAATCG
27301	27401	27501	27601	27701	27801	27901	28001	28101	28201	28301	28501 28601 28701 28801

· 28901 GCCCTACAAC CITGAAGICA GGTILCTIVA ATGITAGGAT CITACITIVAS CCAGGACTIG TCCCGGCGAT TIGTICCAGT CCAACTACAA CCACCACTC

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	CCCATCTIC	COCCATETTO GAACTTCAGT	CCCMAGGALC	TACAGTOCTA	CACTGAAACC	GGINGERTARING			GGT TCATCTC	בול האשלות או	
29001	TANCAGAGAT		ACCAACGGG	בבפכבמנודענ	CCACACTTACA	TUTACCACAA	ATACACCCCA			ACTOGGATAA	
	ATTOTOTOTA	CTGGTTGTGT	TOOTTOCGC	CCCCCCCATTS	GCCTGAATKIT	אהאוזאיזאסא	TATGTCCOOT	TCANACACGG	AAACAGTTAT	すのれたここれです	
29101	CTTCCCCATG	TOGTOGITET	CCATAGCGCT	TATITITIES	TRACTITA	TTATCHERT	CATCTCCTGC	CTARARCCCA	AACOGGCCCCG	ACCACCCATC	
	GAACCCOTAC	ACCACCAAGA	CCTATCCCCA	ATACAMCAT	ACGGAATAAT	NATACACTOA	GTAGACGACG	GATTICOCOT	TOCCCCCCCC	TOCT (CF: FAG	
29201	TATAGTCCCA	TCATTOTCCT	ACACCCAAAC	AATTATTATAA	TUTATAGATT	ההאכמהארדה	AAACACATGT	TOTTITICIES	TACACTATOA	TTANATGAGA	
	ATATCAGGGT	ATATCAGOGT AGTNACACGA	TOTOGGITTO	TTACTACCIT	AGGTATCTAA	CCTGCCTGAC	TITGICTACA	ACAAAAGAGA	ATGTCATACT	AATHTACTCT	
	-1	Xhot									
29301	CATCATTCCT	CATCATTCCT CCAGTITITIA	TATTACTGAC	CCTIGATICC	CTITITING CGINCTECAC		ATTOGCTOCG	OTTICITCACA	TCGAACTAGA	CTCCATTC A	
	GTACTAAGGA	GTACTAAGGA GCTCAAAAT	ATAATGACTG	CCAACAACC	CAAMAMACAC	CAAAAAACAC CCACGAGGTG	TAACCGACGC	CAAAGAGTGT	AGCITICATET	GACOTAAG .T.	
					P _S ()	=======================================					
29401	GCCTTCACAG	OCCUTCACAG TCTATTTGCT	TTACGGATTT	OTCACCCTCA	COCTENTORS CAGGETEATE	CAGCCTCATC	ACTIGICATED ACCEPTITAT		CCAMBCATT	GACTOGGTCT	
	COGAAGTOTC	ACATAAACGA	AATOCCFAM	CACTGCCAGT	GCGAGTAGAC GTCGGAGTAG	GTCCCCACTAG	TCACACCAGT AGCGGAAATA		GCTCACCTAA	CTGACCEANA	
							EcoFl	_\$			
29501	OTOTOCOCIT	TOCATATOTO	AGACACCATC	CCCAGINCAG	GGACAGGACT	ATAGCTGAGC	TTCTTAGAST TCTTTAATTA		TOWNTHAC	TOTOACTT!	
	CACACGCGAA	ACCTATAGAG	TUTOTOTAG		CCTGTCCTGA	TATEGACTEG	ANGMATCTTA AGAMATTAAT		ACTITITAAATO	ACACTGAAAA	
29601	CICCIOATTA	CRECIOAITA TITIOCACCCT	ATCTGCGTTT	TGTTCCCCGA	CCTCCAAGCC	TCAAAGACAT	ATATCATGCA	GATTCACTCG	TATATEGAAT	ATTCCAAGIT	
	DACCACTAAT	BACGACTAAT AAACGTGGGA	TAGACCCANA		CCACCTICUC		TATAGTACGT	CTAAGTGAGC	ATATACCITA	TAAGGITCAA	
							Pstl				
29701	CCTACAATGA	AAAAAGCGAT	CTTTCCCAAG	CCTCGTTATA	TCCAATCATC	TCTGTTATEG	TGPTCTGCAG TACCATCTTA		GCCCTAGCTA	TATATCCCIA	
	COATOTTACT	TITITICGCTA	GAAAGGCTTC	GGACCAATAT	ACCITAGTAG	AGACAATACC	ACAAGACGTC	ATOCTAGAAT	COOGATCCAT	ATATARGUAT	
29801	CCTTGACATT	GOCTOGAACG	CAATAGATGC	CATGAACCAC	CCAACTITICC	כבפכעבבבעפב	TATGCTTCCA	CTCCAACAAG	THOTHOCOGO	COCCUTICN	
	COAACTGTMA	CCGACCTTGC	GITATCTACG	GTACTTGGTG	CONTRANCO	0000000000	ATACGAAGGT	GACGITIOTIC	AACAACGOCC	GCCCANALY 1	
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									Ports		
29901	CCAGCCAATC	Agenteance	Accimented	ACCCCCACTO	AAATCAGCTA	CTTTAATCTA	ACAGGAGAGA ATGACTGACA		CCCTAGATCT AGAAATGGAC	AGAAATGGAC	
	COTCCCTTAG		TGGANGAGG	TOGGGGTGAC	TTTAGTCGAT	GAAATTAGAT	TOTOCHOCHE	TACTGACTGT	GOGATCTAGA	terrracero	
30001	GGAATTATTA	CAGAGCAGCO	CCTCCTAGAA	AGACGCAGGG	CAGCGGCCGA	CCAACAGCGC	ATGNATCAAG			TICCACCAGE	
	CCITAATAAT		GCACCATCTT	1CTOCOTCCC	GTCGCCGGCT	COTTGTCCC	TACTTAGTTC	regreemen	GTACCAATTG	AACTTICOTICA	
30101	GCAAAAGGGG	TATCTTITOF	CTCGTAAAGC	AGGCCANAGE	CACCTACGAC	AGTANTACCA	CCGCACACCG			CCAMOCOT""	
	CGITITICCCC		GAGCATTICG	TCCCOTTTCA	CTCCATCCTC	TCATTATCGT	GECCTOTOGE	GGANTCCATO	TICAACOOTT	GGTTT:CF-AL:T	
30201	GAAATTOOTO	OTCATOCTCG	CACIANAACCC	CATTACCATA	ACTCAGCACT	CRETARAMIC				ACCTGARGAT	
	CTTTANCCAC	CASTACCACC	CTCTTTTCGG	GTAATYSTAT	TGACTCCTGA	OCCARCITIE	GCTTCCGACG	TAAGTGAGTG	DAACAGTTCC	TOGACTOCTA	
				Britis	3						
30301	CTCTCCACCC	CTCTCCACCC TTATTAMERC.	CCTOTOCOOT	CCTATRECAR CICAAAATT TTATTCCCTT	TTATTCCCTT	TAACTAATAA	ANAMANTA	ATTICGING	GAATCAATTT	TAGTCAATCG	
	CAUACUTAGA	GAGACGTGGG AATAATILTG	GENCACECEA	570111	Minner						

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30401	AVATTICITUT	CCAGTETATE	CACCACCACC	AGGAAGGGGA	CCTCCCACACT	CHESTATISC	AGCTTCCTCC	TOCCTOCAAA	CHITCTCCAC	Antetaaatg Tyrgatttae
30501	CAATGICAGE	1100	TCCTGTCCAF	CCGCACCCAC	TATCTFICATE	THEFT	TGAAGCGCGC	AAGACCGTCT	GAACATACCT	TCAACCCCCT
	CITACACTCA		AGGACAGGTA	CARICINATE	ATAGANGTAC	AACAACGTCT	ACTITICATIONS	TTCTGGCAGA	CTTCTATOGA	ACTTRONOC A
30601	GTATCCATAT	GACACOGAAA	CCGCTCCTCC	MCTGTGCCT	THICT TACTO	CTCCCTTTGT	ANCICCCAAT	COSTITICARD	AGAGITCCCCC	TOGGISTACY :
	K141004147		Section 1	Jest Manager I	Par Common	unanananananananananananananananananana				
30701	TETTTACOCC	TATCCGACC	TCTAGTTACC	TCCAATIGCA	TKICTTOCCCT	CAAAATGGGC	AACRONCITOF	CTCTGGACGA	GCCCGCCAAC	CTTACCTCC":
	AGNANCOCOG		ACATCAATGG	ACCTTACCGT	ALGANCOCTA	GTFTTACCCG	TICCCCCCAGA	GAGACCFGCF	CCGGCCGTTG	GANTREACK 1
30801	AAAATOTAAC	CACTIGITORGE	CCACCTCTCA	AAMAACCAA	GTTANATATA	AACCTOGAAA	TATCTOCACC	CCTCACAGTT	ACCTCAGAAG	CCCTANCTAIT
	TITIACATIO	GTOACACTCO	COTOCACACT	THETTHERE	CAGTITICIAL	TTCGACCTTF	ATAGACGICS	OCAGTGTCAA	TOGAGICTIC	GOGNETICACA
10901	OOC TO CCOCC	-	TOGICOCOO	CAACACACTC	ACCATCACAAT	CACAROCCCC	CCTAACCGTG	CACGACTCCA	AACTTAGCAT	TOCCACCCAA
	CCGACGGCGG	CCTCCAGATT	ACCAGCCCCC	CTTCTCTCAG	TESTACCITA	GICTCCGGG	CCATTCCCAC	GIGCIGAGET	TIGARICGIA	Accordage
31001	CONCECCION	CAGTOTCAGA	AGGNANGCTA	COCCHOCAA	CATCAGGCCC	CCTCACCACC	ACCGATAGCA	CATOCCATTAC	TATCACTOCC	TCACCCCC1T AGTCGGGGA1
11101	TAACTE	_	THERECATES	ACTTGAAAGA	CCCCATTTAT	ACACAMANTO	GANAACTAGG	ACTAMAGTAC	OCCUPANT	TOCATIGITAL .
	ATTOATOACO	_	ACCCGTARC	TOANCTITICT	COCCINAATA	TCTCTTTTMC	CTTTTGATCC	TOATITICATO	CCCCGAOGAA	ACCTACATTIVE
31201	AGACCACCTA		CCGTAGCAAC	TOOTCCAGGT	GICACTATITA	ATAATACTTC	CTTGCMACT	ANGITACTO	GAGCCTTOGG	TITICATICA
	TCTGCTGGAT	_	_	ACCAGGTCCA	CACTGATAAT	TATTATGAAG	GAACCITTICA	TTTCAATOAC	CTCGGAACCC	MANCTAN .
31301	CAGGCAATA	TOCANCTIAN	TOTACCAGGA	CCACTAACCA	TREATTETER	MACAGACCC	CTTATACTE	ATOTTACTTA	PCCOPITIOAT	CCTCAAAAC:
	Griccorrat	•	ACATCGTCCT	CCTGATTCCT	NACTAAGAGT	rrrorcrece	CAATATGAAC	TACAATCAAT	AGGCMACTA	CCNOTITION
31401	ACTAMICT	- ANGACTAGGA	CAGGGCCCTC	TTTTTATANA	CTCAGCCCAC	AACTTOCATA	TTAACTACAA	CANAGOCCITY	TACTIONTEA	CAGCTTCAA
	TEATTTAGA	_	GTCCCGGGAG	ANAANTATTT	GAGTEGGGTG	TTGAACCTAT	ANTICATOTT	GTTTCCCGGAA	ATGAACAAAT	GICGRAGITE
		Handia								
31501	CAATTCCAAA		TTAACCTAAG	CACTGCCANG	COCHTCATGE	TTGACCCTAC	AGCCATAGCC	ATTAATGCAG	CACATOGGCT	TONATETEC: F
	THAGGITT	THOOM	AATTCGATTC	GTGACGGTTC	CCCNNCTNCA	AACTOCGATG	TCCCTATCCG	TAATTACGTC	CTCTACCCGA	ACTTANACCA
31601	TCACCTAATO	CACCAAACAC	AAATCCCCTC	AMAACINAAA	TYRICCATOR	CCTAGAATTF	GATTCAAACA	AGCTATOGE	TCCTAAACTA	COMCTRACT
	ACTOCATTAC	: procratoro	TITAGGGGAG	THICHIT	AACCCGGTACC	GGATCTTAAA	CTAACITICT	TCCGATACCA	ACCATITICAT	CCTTGACCCR
31701	TYAGITITION	CACCACAGGT	OCCATTACAG	TAGGNAACAA	MATAATCAT	MOCTAACTT	TETRESACCAC	ACCAGCTCCA	TCTCCTAACT	GTAGACTAAA
1	AATCAAAACT		CGGTAATGTC	ATCCTITICAT	TTTATTACTA	TICGATTGAA	ACACCTRAGIG	TOCTCGAGGT	AGACGATTGA	CATCTCATT
31801	TUCAGAGAAA	. GATGCTAMC	TCACTITIOGS	CTTAACAAA	TUTEXXACTO	MATACTTGC	TACAGITITICA	OFFICER	TTAMOGCAG	THIGGETECA
	ACOTOTOTA		AGTCAAACCA	DAATTOTTT	ACACCGTCAG	TITATGAACG	ATCTCAMOT	CANAACCGNC	AATTICCOTC	AAACCCAGGT
11901	ATAICHERA	CAGITCAAAG	TOCTCATCTT	ATTATAAGAT	TTCACCAAAA	TEXCAGTGCTA	CTANACAATT	CCFTCCFCCA	CCCAGAATAT	T. AACTTA
	TATACACCTT GTCAA	STC.	_	TAATATICTA	ACTOCITIT	ACCTCACGAT	CATTICITA	GOAAGGACCT	OCCUCITATA	ACCTTCAAAT
	2	Byta								
32001	GANATOONCA TCTTM	A TOTTACTON	OCCACANCET	ATACANACOC	TUTTOUNTIL		TATCACCTTA	TCCANANTCT	CACGOTANAN	CTGCCAAAAG
).	CTITACCICI	P AGNATIGACTT	CCGTCTCAGA	TATGITTACG	ACAACCTAAA	TACCICIATTOC	ATAGTCGAAT	ACCITATINGA	GTGCCATTTT	GACGGTTTT

Figure 15T 37/144

32101	TAACATTGTC	AGTICAAGTTT	TGAATTTGCC	ACACAMACT TOTAL TITLES	AAACCTTGTAA	CACTANCOAT TACACTAAAC		CCATGROTCC	AAACAGGGAGA	CACAACTC::A
32201		CINTORCATT			ACAN TACAT		THURCACAT	CCTCTTACAC	TITITICATAC	ATTRICCCON:
32301	ATAAAGAAT	CONTICION	ATCTTTCAAC	CHESTITATE	TTCAATTATA	GAMATTICA	NATIONAL	CATTCAGTAG	TATAGECEEA	CCACCACATA
	TTATTTCTTA	GCAAACACAA		CACAMITAM	AAGTFAACGT	CITTITAAAGIT	TCACTANAA	GTANGTCATC	ATATCGGGGT	GCTGCTCTA
32401	GCTTATACAG	ATCACCGTAC	-	_	CCTAGTATTC		CHICCTCCCA	ACACACAGAG	TACACAGTCC	TTTN: NCCO V:
	CONATATOTC	TACTOCCATO	_	CACTOTOTO	CCATCATAAG	THAGACAGTG	CACOCACCOT	Tereference	ATGTGTCAGG	ANGROSO
32501		AAAAGCATCA	•		TTCTTAGGE	_	CACCGTTTCC	TCTCGAGCCA	AACOCTCATC	ACTONTATT .
		THICGRAGE	_	TICTCTCTAT	ANTANTCCAC	ANTATARGET	GTGCCNAAGG	ACAGCTCCCT	TTCCCAGTAG	TCACTATAL
32601		COOCCAGCTC			ניניאיציאטיאט		TOCTGTCCAA		CTTAACGGGC	GOLDANOTA:
	TATTICAGGG	CCCCGTCGAG	TOANTICARD	TACACCCACA	CELLCUVCCIAC	TCGGTGTCCG	ACGACAGGTT P:st	GAACGCCAAC	GAATTGCCCG	כמיכוווכבוי.
32701	AAGTECACGC	CTACATOGGG	GTAGAGTCAT	ANTCGTGCAT	CAGGATAGGG	CONTROPOS	GCAGCAGCUC	GCGAATAAAC	TECTECCOCC	accacheegr
	Treassioca	DATOTACCCC	CATCTCAGTA	TTAGCACGTA	GTCCTATCCC	GCCACCACGA	conconcoco	CACTTATTTO	ACCIACGGCOG	CGGCGAGGCA
	Path									-
32801	M	TACAACATOO	CACTOCICTC	CTCAGCGATG	ATTICGCACCG	CCCCCCACATCAT	AAROCIACIT		CACAGCAGCG	CACCCITOAT .
	GGACOTCCTT	ATOTTOTACC	Ę	GAGTCGCTAC	TAAGCGTTAT	GGCGTCOTA	TTCCCCCCCAA	CACCACICCC	Grercerece	GTCCCACT**1
			Part							
32901	TCACTTANAT	CAGCACAGTA	~	AGCACCACAA	TATTETACA	AATCCCACAG	TGCAACIGCGC		CCTCATCCCO	OCCACCACAG
	AGTCAATTTA	Greenoreat	TCACCTCCTG	TCCTOGTGTT	ATANCMOTT	TTAGGGTGTC	ACGITICCIACO	ACATAGGITT	CONGINCOCC	cccromanc
33001	AACCCACOTO	GCCATCATAC	CACAAGCGCA	CCTACATTAA	GTRICCGALCC	CTCATABACA	CGCTGGACAT	AAACATTACC	TCTTTTGGCA	TCTTGTAATT
	THOOOTIGEAC	COGTAGTATO	GICTICGCGT	CCATCTAATT	CACCGCTGGG	GAGTATTIGT	CCCACCTGTA	TITICTAATOG	AGNARACCOT	ACAACATTAA
		Kpri								Ped ·
33101	CACCACCTCC	COCTACCATA	TAAACCTCTG	ATTARACATO	GCGCCATCCA	CCACCATCCT		GCCAAAACCT OCCCGCCGGC	acceaceace	TATACACTIX.
	GTGGTGGAGG	GCCATGGTAT	ATTROCACAC	TAATTIGIAC	CGCGSTAGGT	GOTTOTACCA	THICKNOOP	TGCA	دومودوودده	ATATIOTO:
								Ecolly		
33201	ACCOMACCOG CACTOGAACA	GACTOGAACA	ATGACAGING	AGAGCCCAGG	ACTOTTAACC	ATCICATICATIC	ATCCTCGTCA	TCATATCAST (GTTGGCACAA	CACAGGCACA
	TCCCTTGGCC	CTOACCTTOT	TACTUTCACC	TCTCGGGTCC	TCACCATTOS	TACCTAGTAG	TACCAGCAGT	ACTATAGTTA	CAACCGIOIT	Grenticenci
	_									Pell
33301	COTOCATACA	CTTCCTCAGG	ATTACARACT	CCTCCCGCnT	TACAACCATA	TCCCAGGGAA	CAACCCATTC		GTAAATCCCA	CACTGCAGGG
	OCACOTATOT	DANGGAGTCC	TAATGTTCGA	GGANDCACOCA	ATCTINGTAT	AGGETCCCTT	GETGGGTANG		CATTTAGGGT	GTGACCTCC
33401	MONCETCOC	ACCITAACTCA	COTTGTGCAT	TOTOMACTO	TTACATTCOS	CHINCAGGGT			CCCCCCTTTC	TOTOTOWN
	TTCTOGAGCG	TOCATTICAGE	CCANCACGTA	ACAGITATOAC	ANTOTANGEC	נייובייוביייב	TACTAGASACIO	TCATACCATC	OCCICCOAAA	ACAGAGETET F
33501		GATCCETACT		-	ACCGAGATCG	Test Tests for			OCCOCACOTA SCOTTOCAR	CTCATATTT .
	CCTCCATCTO	CTAGGGATGA	CATACCTCAC	accocneror	TOCCICIAGE	ACMACCAIACA	TCACAGTACG	GITIMCCITO	COCCIOCAT	CASTALANS

ACATEGRAGIT GTAGTATATE CACTETETEA ACATEACAC CATEGRAFIT CACCACEGCA GAATAAGECA CACCECAGCC/ GTGGTGGCGT CTTATTCGGT GTGGGTCGTT TTATTCCAAA AGATTATCCA AAACCTTAAA	ATCOCATTTO TANGATGTTO CACINITOGCT TACCGTAAAC ATTCTACAAC GTOTTIACCG/ TAAACAFTCC ACCACCTTCA ACCATGCCCA ATTTOTAAGG TCOTGAAGT TCGTACGGT	AAAANCTGC TCCAGAGGG CCTCCACTT TTTTAGACG AGGTCGCG GGAGGTGGAA GAACATTAAC AAAAATACCG CGATCCCGTA CTTGTAATTG TTTTTATGGC GCTAGGGCAT	ACCTICATE ACAAAGAAC CCACACTGAT ACCTICATAC TOTITICITO GGTGTGACTA ATAAAATGCA AGGTGCTOCT CAAAAATCA	TCCACGACGA AGAAAAAGAC TCTTTTTCTG	CTCCTCGGTC GAGGAGCCAG	TATGTATGGGA GCCTAGGCAA CGGATCGGTT	THEOGRAM THEFFERE CHARCETORS THEOGRAM GITAMBEC ACMANAGA ACTOCCATIC CARTIFCAGE TOTITITIS ACTICCETT TOCCACETTA COTCACTTOC TOMOGENA AGGEGENT GENTERAGE
GATCCCTCTG CTATCCAGAG TYATAACATC ACTATTCTTT GTTTTTTTT CAANAMAAAA	AGAACAGATA TCTTGTCTAT ATCTCCTCTA	CGGCCATTGT GCCGGTAACA TTCAAAAGCG	CTTCCCCGCC	CCCCCCCCTA GTAAGCTCCG CATTCGAGGC	ANCATTAGAA TTGTAATCTT TTAAAAAGCA AATTTTTCGT	OACCOMATA CTGGCTTTAT AACACCTGAA TTGTGGACTT	AGTANANAAG TCATTTTTC GACTANAAA CTCATTTTTT TCAAATCGTC AGTTTAGCAG
GTCTTCGGTC TYTCGGCTTA CAUAGGCCAAT TCCTTCATGG GCTCCTTCCC AGGAAGTACG CGCGACGGG GGAAGAGCTG GAAGAACCAT CCTTCTCGAC CTTCTTGGTA	TRIGICIANCT CTACAGCCAA ACCAGTITICA GATGICGGTT GGCTAAAACCC TICAGGIGA		TGCACGGACC AGCGCGGGCATCA ACGTGCCTGG TCGCGCCGGT AGGGCCGGT AGGCCGGT AGGGCCGGT AGGGCCGGCGCGT AGGGCCGGCGT AGGGCCGGT AGGGCCGGCGT AGGGCCGGT AGGGCCGGCGGT AGGGCCGGT AGGGCCGGT AGGGCCGGCGCGGT AGGGCCGGCGCGGT AGGGCCGGCGCGGT AGGGCCGGCGGT AGGGCCGGCGGT AGGGCCGGCGCGGT AGGGCCGGCGCGGT AGGGCCGGCGCGGT AGGGCCGGCGGT AGGGCCGGCGGT AGGGCCGGCGCGGT AGGGCCGGCGGT AGGGCCGGCGCGGT AGGGCCGGCGGT AGGGCCGGCGGT AGGGCCGGCGGT AGGGCCGGCGCGGT AGGGCCGGCGCGGT AGGGCCGCGGT AGGGCCGGCGGT AGGGCGCGCGGCGCG			ATCGGTCAGT GCTAAAAAGG TAGCCAGTCA CGATTTTTCG ATAGGAGAGA AAAACACATA TATCCTCTCT TTTTGTGTAT	CCATANCAGT CAGICTTAICE GGTATTGTCA GTIGGAATGG TGCAGAGCGA GTATATATATA ACGTCTCGCT CATATATATG CCAANAMACC CAGAACTTCC GGTTTTTTGG GTGTTGAAGG
Hgitt ACA'ATCTAGACG GTATGTAAAG GATACATTTG GGGAAAJAGCG CCCTCCTCCC	TCCGGTGGCG AGGCCACGG TCGACGTAAA	TATCTCTAAG ATAGAGATTC AATTCAGGTT	CGACGTCCAG	CAGCGTAGCC GTCGCATCGG TCGTAGTCAT AGCATCAGTA	ACACAAAATA TGTGTFTTTAT GCGTGACCGT CGCACTGGCA	GINCHANGED CANCEANGED AACAAMITA FIGHTENAE	ACAGGGGAA HGTCGCCGTC AAGGGCCAAG TTCCCGGTTC GAAACGAAAG
TGCG GGCGTGACAA ACGC CCGCATTGTT CCTG GCTTCGGGTT GGAC CGAAGCCCAA TGCG AGTCACACAC		CCAC CITCICMIN COTO GAGAGITAT UTCA TGATICCAAA	• • •	COMO CTATOCTANC SCCTC GATACGATTG WAAA AGAANGCACA FTTTT TCTTTCGTGT	COCCA ANGACGIANT ACTAC COCCATOCCG IGATO CCOCTACCGC	COCAT TROTOTAGE CCCCA TROTOTAGE CCCCA TROUBURTATE CCCCA TROUBURTATE CCCCATA	NGATA CAGCGCTTCC TOTAT GTCCCGAAGG NGTCA CAGTGTAAAA TCAGT GTCACATTTT GCGAA CCTACGCCCA CCCTT GGATGCGCCT
CTGAAGCAAA ACCAAGTGCG GACTTCGTTT TGGTCCACGC AAGCATCCAG GCGCCCCTG TTCGTAGGTC CGCGGGGAC ACCTACACAT TCGTTCTGCG	\$			FATGACACC ATACTCGGAG ATACTGGCC TATGAGCCTC GACAAAGCCT CGCGCAAAAA CCGTTTCGGA GCGCGTTTTT	TCTCAACAT OTCTCCOOPT AGACTITOTA CAGACOCCA ATAACCATAA GACOGACTAC TATTCGTATT CTGCCTGATO	TCATAATOTA AGACTCOOTA AGTATTACAT TCTGAGCCCAT AGACAACATT ACAGCCCCCA TCTGTGTAA TGTCGGGGGT	
33701 A	33901	34101		34401	34601	34901	35001 35101 35201

Figure 15V

PMRKANISGOG MER6A2

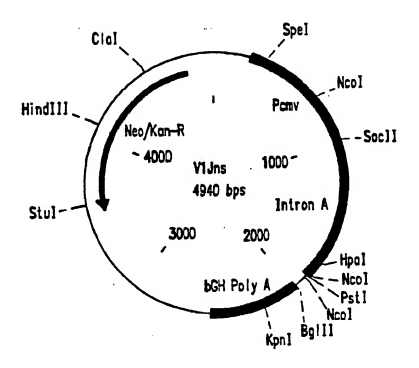
TITICATOTTA ACCOTTICACI ATCITICAATG ACCOTICAT TICGATOCIAS TORCOCCOCC AACCOTOCICO GOCCOCOTOC ACTITITAAG CTACCOOLAG GICCOMOTTIC TCACAMOTO CGACTTCACT CACCCCTCA TTATCATATE GOTTICAATE CAAAATAAGG TATATTATTG ATCATGTAA TTANGAATTE GGATCTGCGA CGCRAGGCTG GATGGCCTT **GCTCAAGTC** TEGAGGAGE CAGGAGGETA GEMENTIET TEATTENNEE GOEGICACAA TAGGAGTAE CAATACEGTE GTGACGTAIT AAGAGAATEA ATOCCCTAT AGTATTICO TCATAMCC AACAAACT GTTANGGGA GTTINCONATI CGNATCKITAC GICTICACCA ACTAGGGGGT ACAACACGTT OCCATICTAC GAAAAGACAC TOACCACTCA TOAGITGGIT CAGTAAGACT CITATCACAT ACGCCGCTGG CTCAACGAGA ACGGGCGA CTROCCTO GACCCGACA TOOCAGCAG THEFTINE CAATTCCCT/ **GCTTACCATA** ACCENCENC CCCCCCACG AATAGTATAA CCGAAGTTAG GTTTTATTCC ATATAATAAG TACTACAATT AATTCTTAAG CCTAGACGCT GCGCTCCGAG THUCOGGGG ATCGCATHE CONCEPTED ASCANGEN TUCAGEDG FAGATIACOA CCATCAGGA OCTAGTCCCT ANAMATICOAC OCTOGTAGTG TTTTTAGCTG OCCACCAMA TECHTICATE TITIETACGO GOTETGACGE TEAGTGGAAC GAANACTEAC CCTGTTCCGA CCCTGCCGCT **GCITACGGCGA** TCCCTCCAAG ACCOAGGITC TTATCOCCAC **AATAGCCCTC** CTAGANGCAC DATICTICCTO COCTOCTTT CTTTTCAGTO TOACAGTTAC ACTOTICAATO ATACOGGAGO TATOCCCTCC **GCGCAACGTT** ACCICCTICG GICCICCGAT COTHCINCAGA AGTANGITUG CCGCAGIGIT ATCACICATO GITATOGGAG CACTGCATAA CCUTANGAIG CITITICITIS ACTIVITIENT ACTIVANCEAN GICATICIDA GAATAGIGIA FORGOCGACE GAITITICIO CCCCCCTCCC COCOTTOCAA TACAAGTTAC TYCORCCTAA AACCTACGTC ACCGGCCCCG TYCCCAGGCC GOCOCIANCOT CCASTACGAC AGGICCOTCC ATCTACTGCT TATCHARTIC GACATECATA GALTICAARCE ACATECAGCA GCAGAACTCA GCITCGGCCA TTCTGTGCTG TUTTICANGTO GTOGCCTAAC TACOGCTACA CTCGCTCCAT ACATCCCCCA CGATCTCTCA AGAACTTCAC CACCGGATTG ATGCCGATGT AGITACCITIC GGAAAAAGAG TITXJIMACITC TITATCCGGC AAACAAACCA CCCCTGGING TITITICCIA GAGIICTICI AGGANACIAG ANAGANICC CCAGACITAG AGICACCITO THICCINGAL GREGARCIAG GAAAATITAG TIAGATITICA TATATACICA TITGAACCAG CONCCATCAC TOTAGGTCOF **GOCGACCATIC** CCACCCCCAA GTAGGTCAGA TAATTAACAA CGGCCCTTCG ATCTCATTCA TCAAGCGTC AATTATCAAA PRECTAGTIC COCTCAATOT CCACAAGGCT AAGACACGAC AAACTIKGGIC GATAACTACG CTATTICATION GOTCOCCTT TTAATAGTTT GCCAOTTACA AGATTATICAN, AANGGATCTT CACCTAGATC CTTTTAAATC AATCTAANGT ATATATGAGT GCATTITICE GOEGENACIGA CEGENANANG GTATECICIAGE COOCCICAET GOGDAGCHIG GCGCTTTCTC ATARCICACG CTGTACGTAT CTCANTICGG CCTTTTTCTC NACCATCGAG AACTAGGCCG TITGTTTGGT COCTAGACAG ATAAAGCAAG TAGGTATCAA CRGACTGAGG GGCAGCACAT PACCOCCAGA CCCACGCTCA CCGGCTCCAG ATTATATAGACCAG GCCGGGAAGC TAGAGTAAGT AGTTCGCCAG COCICGICGE FIGGRANGE TECATICAGE TECNSTREEC AAGANTEAAD CATAGRETICE GEOECECTICA TITICCCCTIC GAAGCIICCCT CGIGGGCTCT AANGRIGGGAC CTTCGAGGGA GCACGCGAGA CCANCCCOOF OCCTOACTCC CCOTCOTOTA TATTEGET OCCARCAGIA ANCIATACIC ANGTANGTICO AGGICANGGO COGTAACTAT COTCTTGAGT TANATAGECO CCCSTTTTC **GCCATTIGATA** CCTACACAGT ATCCATAGIT ATOCCOCTCT GOCTOCCAGT CCCCCAGGTC TAGCCCTACG GACCGAGGTA TOTAGGCGGT ANANANGGAT CTCAAGAAGA GACTATAAAG NTACCAGGCO ANGANGGANG CCCTTCGCAC CCCGANACAG COCOGNATAG CCCCCTTCCT TATAGREECOC CCCGACCGCT GCGCCTTATC TATTACGITIC CATCCAGTCT ATTAATTGTT TCCCAACACA AAGGCCGCCG CCTTAAAAAGG CTGATATTTC OGGCTGGCGA *fcantidand* OCCURACTOR AAACTACAAT TUCCOCTIC CCCCOTTCAD THEFTETEGE AAGAAGAGC GOCCAGGNAC CCOCICCTIG AACCCCAACAG TIGGGCTGTC COCCAAGIC CACTOSTANC AGGATTAGCA TCCTAATCOT TOCTONAGEC ACGACTTCGG TACGCGCAGA ICOTOGRETA ATGCGCOTCT TCTAATAGTT TOGATAGAGT **GCTGCAATGA** COACGITACT PATECOCCIC ATAGGCGGAG **GCACCACAGT** ACCTATCTCA COTOCHURCA CTOGGGGAGT **GCOTANTACT** CACACOCOCA DTUACCATTO THOOTCATO TTTTCCCCAA GTCATGCCAT CAUTACOUTA CATTTTANGA **GTAMATTIC!** CCCATTATOA COOLCONTIN DACOTOCCOA CTCCACCGCT CTUTCCOCCT **ICCTIOCTIOD** ATCTOCCCTC PADACGCCAG ICCACCACAT MACCAGTAC **IGTICACTICCO ACCOSOSICA** DATOTCCOTA AAAACCOGTT CCCACCAMA TOCACOAACC CAOTOAGGC TOCCCCAOT CCTCCACTT CACOTTOAA CTACAGGCAT 35301 35401 35601 35701 35801 36001 36101 36201 36301 36401 36501 10991 36801 36901 35501 35901 36701

figure 15W

PMRKAd5gag MER682

37001	CAACACGODA	CANCAGGODA THATAGGGGG CCACATARGA GANCTITANA AGTRITISTI ATTRIANAAG GITCITIGGG GGGAAAGTE ICAAGGAICT TAGRICITATT	CCACATARCA	CANCTITANA	AGTYXCTYCATY	ATTRICAMMAC	GENCHICOGG	CCGNNACTC	TCANGGATCT	TACCHETA
	Gritchaccet	OFFORECCT AFFANCECE COTOFATER CFICALAFFF TCACACTAR TAACCTFFTG CAACAAGICC COCFFFICAG AGFOCFAGA AFGICAAAAA	OCTOTATEGE	CHICAANITI	TYNACANGTAG	TAACCITITIG	כאשנישיטעיכנ	CCCTTTTCAG	NGPTCCTNGA	ATGGCGACAA
37101	GAGATCCAGT	ONDANCEACT TEGATOTAC CENETECING ACCENATION TETRINISTAT CITITACTIT ENCONSCIENT TETRICAGING CARABACHES ANGENAAT	CCACTCGTGC	ACTENACTES	TKTTY:AKTAT	CTITITACTIT	CACCAGGGT	TCTCCCTCAG	CAANAACACO	AACCCAAAAT
37201	GCCGCAAAA	CICINGOLA ACCIACATIO CITTATATO INTERIORA TACTOTATATO CITTOTOTO CITATATO CANGENTIA TCATATATA TOTOTOTATA TOTOTOTATA TCATATATA TCATATATAT	OCCUPACACO:	ANATIGITICAN	TACTICATACT	CHICCHITT	CANTAITATT	GANGCATTTA	TCAGAGITAT	TCTCTCATCA
37301	GCGGATACAT	CCCCATCA ATTICAATOR ATTACAAA ATAACAAA AGGATTAC GATAAAAA GAGATAAAA AGGATAAAAAAA GAGATAAAAAA GAGAAAAAAAA	ATTTAGAMAA	ATAMACAMAT	AGGICA:TTTCTC	CCTCTATTTC CCCTCTAAAG	CCCGAAAAGT	COCTEGACTE	GTCTAAGAAA	CCATTATTA
						- Ban	F.colli			
37401		CATGACATTA ACCTATABADA ATAGGGGTAT GAGGAGGCC TTICGTCTC ANGANTIGGA TIGGAATTCT TAAT (SEQ ID NO: 27) GTACTOTABAT TOGATATTTT TATCCGCATA GTGCTCCTGG AAAGGAGAAG TTCTTAACCT AGGCTTAAGA ATTA (SEQ ID NO: 28)	ATACCCCATA TATCCCCATA	CACGAGGCCC	TTICGTCTTC MAGCAGAG	AAGAATITGGA TTCTTFAAGGGT	TUCGAATTET ACCETTAAGA	TRAT (SEQ ATTA (SEQ	ID NO: 27) ID NO: 28)	

Figure 15X



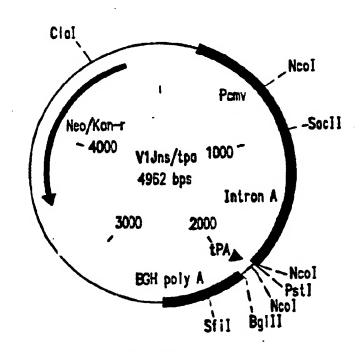


FIGURE 16

GCACTGGCCCCTGACTGAGGAGAGATCAAGGCCCTGGTGGAAATCTGCACTGAGATGGAGAAGGAGGGGCAAAATCTCCA sGInTrpProLeuThrG1uG1uLys11eLysA1uLeuVo1G1uI1eCysThrG1uMelG1uLysG1uG1yLysI1eSerL 30 40 50

AGATTGGCCCCGAGAACCCCTACAACACCCCTGTGTTTGCCATCAAGAAGAAGAAGAACGACTCCACCAAGTGGAGGAAGCTGGTG
ysleGlyProGluAsnProTyrAsnThrProVolPheAlolleLysLysLysAspSerThrLysTrpArgLysLeuVol
60 70

GACTICAGGGAGCTGAACAAGAGGACCCAGGACTTCTGGGAGGTGCAGCTGGGCATCCCCCACCCCGCTGGCCTGAAGAA AspPheArgGluLeuAsnLysArgThrGlnAspPheTrpGluVolGlnLeuGly1leProHisProAloGlyLeuLysLy 80 90 100

GAASAASTCIGTGACIGTGCTGGCTGTGGGGGATGCCTACTICTCTGTGCCCCTGGATGAGGACTTCAGGAAGTACACTG slyslysSerVolThrVolLeuAloVolGlyAspAloTyrPheSerVolProLeuAspGluAspPheArgLysTyrThrA 110 120 130

CCTTCACCATCCCTCCATCAACAATGAGACCCCTGGCATCAGGTACCAGTACAATGTGCTGCCCCAGGGCTGGAAGGGC
InPheTnrlieProSerlieAsnAsnGluThrProGlylleArgTyrGlnTyrAsnVolLeuProGlnGlyTrpLysGly
140 150

TCCCCTGCCATCTTCCAGTCCTCCATGACCAAGATCCTGGAGCCCTTCAGGAAGCAGAACCCTGACATIGTGATCTACCA SerProAlollePheGinSerSerMetThrLyslieLeuGluProPheArgLysGinAsnProAsplieVollleTyrGI 160 170 180

CTACATGGCTGCCCTGTATGTGGGCTCTGACCTGGAGATTGGGCAGCACAGACCAAGATTGAGGAGCTGAGGCAGCACCACC
nTyrNetAioAtoLeuTyrVotGtySerAspLeuGtuIteGtyGfnHisArgThrLysIteGtuGtuLeuArgGfnHisL
190 200 210

TGCTGAGGTGGGGCCTGACCACCCCTGACAAGAAGCACCAGAAGGAGCCCCCCTTCCTGTGGATGGGCTATGAGCTGCAC euLeuArgTrpGlyLeuThrThrProAsplysLysHisGlnLysGluProProPheLeuTrpMetGlyTyrGluLeuHis 220 230

CCCGACAGTGGACTGTGCACCCCATTGTGCTGCCTGGAGAAGGACTCCTGGACTGTGAATGACATCCAGAAGCTGGTGGG ProAsplysTrpThrVoIGinProIieVoILeuProGluLysAspSerTrpThrVoIAsnAspIieGinLysLeuVoIGI 240 250 260

CAAGCTGAACTGGGCCTCCCAAATCTACCCTGGCATCAAGGTGAGGCAGCTGTGCAAGCTGCTGAGGGGCACCAAGGCCC yLysleuAsnTrpAloSerGin]leTyrProGiyileLysVolArgGinLeuCysLysleuLeuArgGiyThrLysAloL 270 280 290

FIGURE 17A

GCCCTGTACTATGACCCCTCCAAGGACCTGATTGCTGAGATCCAGAAGCAGGGCCAGGGCCAGTGGACCTACCAAATCTA GiyVoITyrTyrAspProSerLysAspLeulieAloGiulieGlnLysGlnGlyGlnGlyGlnTrpThrTyrGlnIleTy 320 330 340

CCAGGAGCCCTTCAAGAACCTGAAGACTGCCAAGTATGCCAGGATGAGGGGGGGCCCCACACCAATGATGTGAAGCAGCTGA rGInGIuProPheLysAsnLeuLysThrGIyLysTyrAIoArgMeLArgGIyAIoHisThrAsnAspVoiLysGInLeuT 350 360 370

CTCAGGCTGTGCAGAAGATCACCACTGAGTCCATIGTGATCTGGGGCAAGACCCCCAAGTTCAAGCTGCCCATCCAGAAG hrGiuaioVoiGinLyslieThrThrGiuSerlieVoilieTrpGlyLysThrProLysPheLysLeuProlleGinLys 380 390

GCTGAAGCTGTGGTACCAGCTGGAGAAGGAGCCCCATTGTGGGGGGCTGAGACCTTCTATGTGGCTGGGGCTGCCAACAGGG uVoilysLeuTrpTyrGinLeuGiuLysGiuProileVoiGlyAloGluThrPheTyrVoiAioGlyAloAloAsnArgG 450

AAGACTGCCCTCCAGGCCATCTACCTGGCCCTCCAGGACTCTGGCCTGGAGGTGAACATTGTGACTGCCTCCCAGTATGC LysThrAloLeuGInAlolleTyrLeuAloLeuGInAspSerGiyLeuGiuVolAsnileVolThrAloSerGInTyrAl 480 490 500

CCTGGGCATCATCCAGGCCCAGCCTGATCAGTCTGAGTCTGAGCTGGTGAACCAGATCATTGAGCAGCTGATCAAGAAGG aleuGlylielleGlnAloGinProAspGInSerGluSerGluLeuVolAsnGinIielleGluGinLeuIleLysLysG 510 520 530

AGAAGGTGTACCTGGCCTGCCCGCCCACAAGGCCATTGGGGGCAATGAGCAGGTGGACAAGCTGGTGTCTGCTGGC
IULysVolTyrLeuAloTrpVolProAlgHisLysGlyIleGlyGlyAsnGluGInVolAspLysLeuVolSerAlgGly
540
550

ATCAGGAAGGTGCTGTTCCTGGATGGCATTGACAAGGCCCCAGGATGAGCATGAGAAGTACCACTCCAACTGGAGGGCTAT

11eArgLysVolleuPheleuAspGlyi1eAspLysAloGinAspGluHisGluLysTyrHisSerAsnTrpArgAloMe
560 570 580

FIGURE 17B

GGCCTCTGACTTCAACCTGCCCCCTGTGGTGGCTAAGGAGATTGTGCCCTCCTGTGACAAGTGCCAGCTGAAGGGGGAGG tAloSerAspPheAsnLeuProProVolVolAloLysGlulleVolAloSerCysAspLysCysGlnLeuLysGlyGluA 590 600 610

GCTGTGCATGTGGCCTCCCGCTACATTGAGGCTGAGGTGATCCCTGCTGAGACAGGCCAGGAGACTGCCTACTTCCTGCT AlovalHisValAloSerGlyTyrlleGluAloGluVallleProAloGluThrGlyGlnGluThrAloTyrPheLeuLe 640 650 660

GAACCTGGCTGGCAGGTGGCCTGTGAAGACCATCCACACTGCCAATGGCTCCAACTTCACTGGGGCCACAGTGAGGGCTG
uLysLeuAloGlyArgTrpProVolLysThrlleHisThrAloAsnGlySerAsnPheThrGlyAloThrVolArgAloA
680
690

CCTGCTGGTGGGCTGGCATCAAGCAGGAGTTTGGCATCCCCTACAACCCCCAGTCCCAGGGGGTGGTGGCCTCCATGAAC
IOCysTrpTrpAioGlylleLysGInGluPheGlylleProTyrAsnProGInSerGInGlyVolValAIoSerMelAsn
700 710

AAGGAGCTGAAGAAGATCATTGGGCAGGTGAGGGACCAGGCTGAGCACCTGAAGACAGCTGTGCAGATGGCTGTGTTCAT LysGluLeuLysLyslielleGlyGlnVolArgAspGlnAloGluHisLeuLysThrAloVolGlnMeiAloVolPhell 720 730 740

CCACAACTICAAGAGGAAGDGGGGCATCGGGGGGCTACTCCGCTGGGGAGAGGATTGTGGACATCATTGCCACAGACATCC
eHisAsnPhelysArglysGlyGlylleGlyGlyTyrSerAloGlyGluArglleVolAsplleIleAloThrAsplleG
750
760
770

AGACCAAGGAGCTCCAGAAGCAGATCACCAAGATCCAGAACTTCAGGGTGTACTACAGGGACTCCAGGAACCCCCTGTGG
InThrLysGIuLeuGInLysGin!IeThrLysIieGInAsnPheArgVoITyrTyrArgAspSerArgAsnProLeuTrp
780
790

AAAGCCCGGGCAGATC" (SEQ ID NO: 3)

Xx Bg/l1 (SEQ ID NO: 4)

FIGURE 17C

(within SEQ 10 NO: 7) CCACCCACATOTOCOCCCATOTOCCCATTOACACTOTOCCTGTCAAGCTGAACCCTCCCATGCATCCC RoserGiulleSerAidProlieSerProlieGiuThrVoiProVoiLysLeuLysProGlyMeiAspGly 20 20 20

FIGURE 18

WT OPT	- ATG GGT GGC AAG TGG TCA AAA CGT AGT GTG CCT GGA TGG TCT -4	42
OF 1		14
WT		84
OPT	- ÁCC GTG ÁGG GÁG ÁGG ÁTG ÁGG AGG GCC GÁG CCC GCC GÁC T V R E R M R R A E P A A D -:	28
ਘਾ	- AGG GTG AGA CGA ACT GAG CCA GCA GCA GTA GGG GTG GGA GCA	126
DPT	- AGG GTG AGG AGG ACC GAG CCC GCC GCC GTG GGC GTG GGC GCC	42
WT	- GTA TCT CGA GAC CTG GAA AAA CAT GGA GCA ATC ACA AGT AGC	168
OPT	. GTG TCC AGG GAC CTG GAG AAG CAC GGC GCC ATC ACC TCC TCC	-56
WT		210
OPT	. AAC ACC GCC GCC ACC AAC GCC GAC TGC GCC TGG CTG GAG GCC	-70.
WT .	- CAA GAG GAT GAG GAA GTG GGT TTT CCA GTC AGA CCT CAG GTA	-252
OPT	· CÁG GÁG GÁC GÁG GÁG GTG GGC TTC CCC GTG AGG CCC CAG GTG	-84
WT	- CCT TTA AGA CCA ATG ACT TAC AAG GGA GCT GTA GAT CTT AGC	-294
OPT	- CCC CTG AGG CCC ATG ACC TAC AAG GGC GCC GTG GAC CTG TCC	-98
WT	- CAC TIT TTA AAA GAA AAG GGG GGA CTG GAA GGG CTA ATT CAC	-336
OPT	- CAC TTC CTG AAG GAG AAG GGC GGC CTG GAG GGC CTG ATC CAC	-112
₩T	- TCA CAG AAA AGA CAA GAT ATC CTT GAT CTG TGG GTC TAC CAC	-378
OPT	- TCC CAG AAG AGG CAG GAC ATC CTG GAC CTG TGG GTG TAC CAC SQKRQDILDLWVYH	-126
WT	- ACA CAA GGC TAC TTC CCT GAT TGG CAG AAC TAC ACA CCA GGG	-420
OPT	- ACC CAG GGC TAC TTC CCC GAC TGG CAG AAC TAC ACC CCC GGC T Q G Y F P D W Q N Y T P G	-140

FIGURE 19A

WT OPT	II II III II II II II II II II II III III III III III III III CCC GGC ATC AGG TTC CCC CTG ACC TTC GGC TGG TGC TTC AAG	462
		154
₩T	11 11 11 11 11 11 11 11 11 11 11 11 11	504
OPT	CTG GTG CCC GTG GAG CCC GAG AAG GTG GAG GAG GCC AAC GAG L V P V E P E K V E E A N E	- 168
WT	GGA GAG AAC AAC TGC TTG TTA CAC CCT ATG AGC CAG CAT GGG	- 546
OPT	GGC GAG AAC AAC TGC CTG CTG CAC CCC ATG TCC CAG CAC GGC	-182
WT	All Blo Glo Glo Glo Glo Glo Glo Glo Glo Glo G	-588
OPT	- ATC GAG GAC CCC GAG AAG GAG GTG CTG GAG TGG AGG TTC GAC	-196
WT .	- AGC AAG CTA GCA TTT CAT CAC GTG GCC CGA GAG CTG CAT CCG	-630
OPT	- TCC AAG CTG GCC TTC CAC CAC GTG GCC AGG GAG CTG CAC CCC S K L A F H H V A R E L H P	-210
WT	- GAG TAC TAC AAG GAC TGC TGA (SEQ ID NO:30)	-651
OPT	- GAG TAC TAC AAG GAC TGC TAA (contained within SEQIDNO:9) E Y Y K D C (SEQIDNO:10)	-216

FIGURE 19B

VIJIIS/IIIE) *Pst! Bg1II* Cataggictiti<u>ciaca</u>gicaccotcctta<u>ggaic</u>iaccacc atg ggc ggc agg tag tag tag agg tag tag ggc . M G G K W S K R S V P

Srf1 Bg111

. . . . CAC CCC GAG TAC TAC AAG GAC TGC TAA AGCCCGGGCAGATCTGCTGCCTTCTAGTTGCCAGC (SEQ 1D NO: 38)

H P E Y Y K D C * (contained within SEQ 1D NO: 10:

VIJns/nef(G2A.LLAA)

Psti Bgili Catrgrettticiacagicaccgiccttgagaictgccacc atg gcc ggc ang tgg tcc aag agg tcc gtg ccc . M A G K W S K R S V P

SrfI BOILI . . . CAC CCC GAG TAC TAC AAG GAC TGC TAA AGCCCGGGCAGAICIGCCTGTGCCTTCTAGTTGCCAGC (SEQ ID NO: 39) H P E Y Y K D C * (contained within SEQ ID NO:14)

V1Jns/tpanef & V1Jns/tpanef(LLAA)

CATIGGETETTTTCTGCAGCCTCTTATATCTAGATCACC ATG GAT GCA ATG ANG AGA GGG CTC TGT 1GT GTG N N N N N G L C C V

CTG CTG CTG TGT GGA GCA GTC TTC GTT TCG CCC AGC $\frac{BgIII}{S}$ CC TCC AAG AGG TCC GTG CCC . . .

SrfI Bg111

CAC CCC GAG TAC TAC AAG GAC TGC TAA AGCCCGGGCAGAICIGCTGTGCCTTCTAGTTGCCAGC (SEQ 1D ND: 40)

H P E Y Y K D C * (contained withon SEQ ID NO: 16)

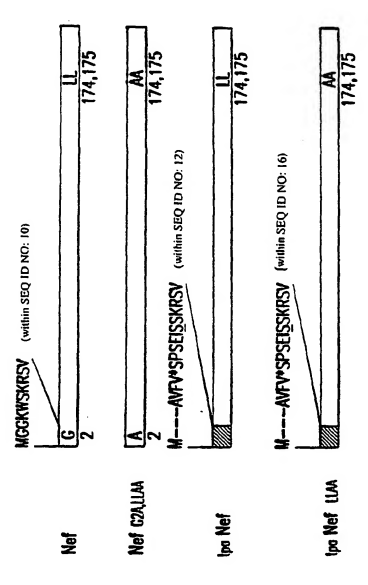


FIGURE 21

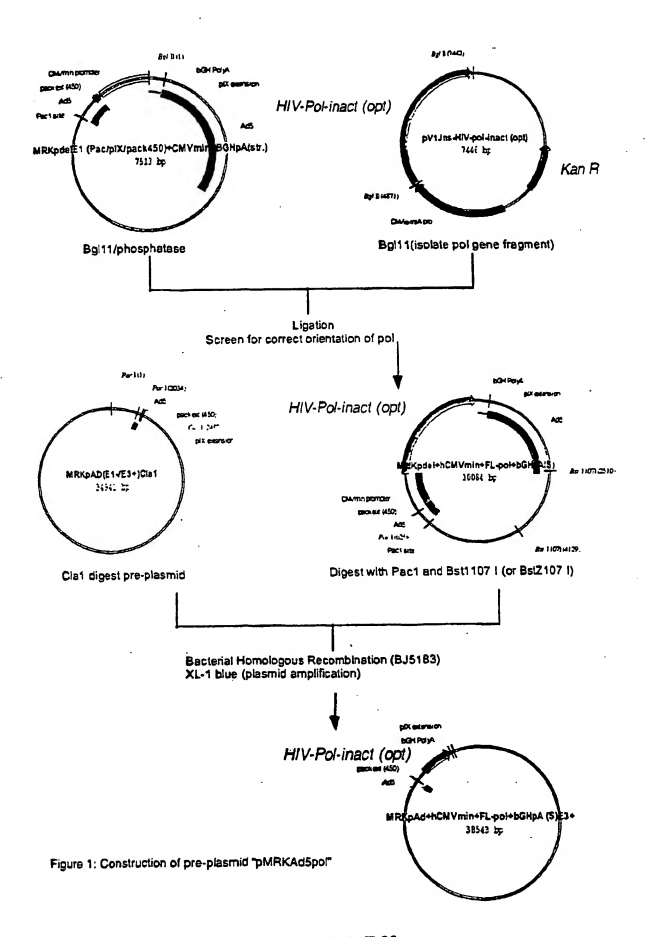
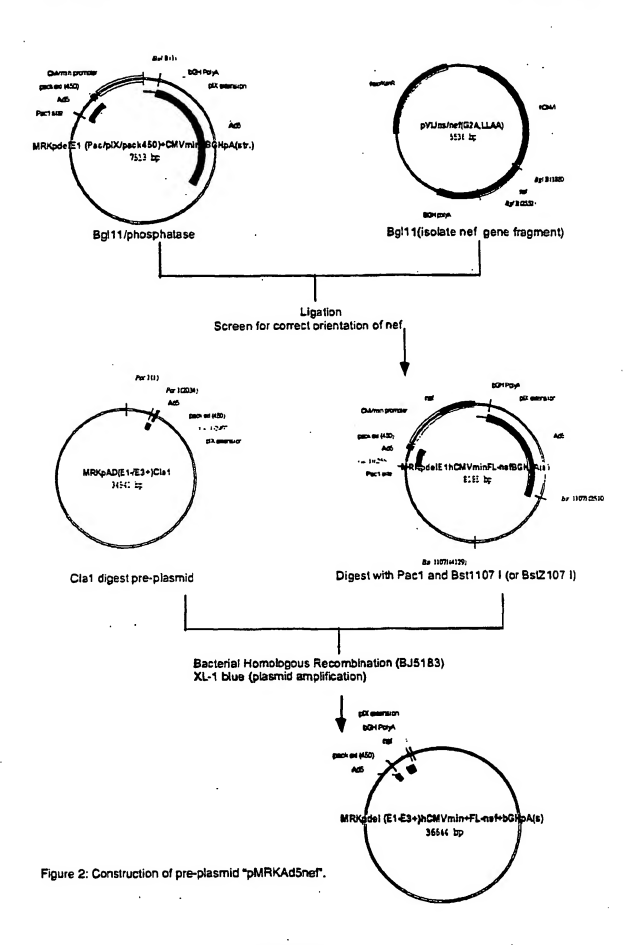
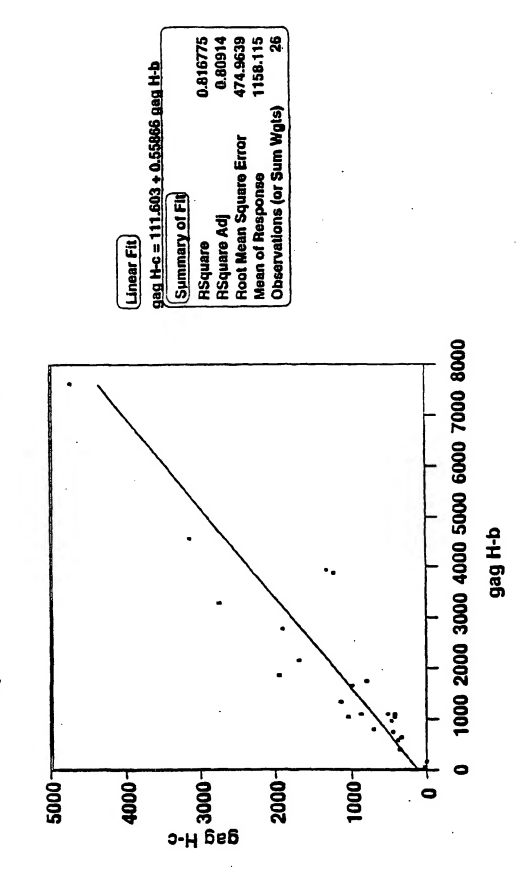


FIGURE 22



Comparison of Clade B vs. Clade C Anti-gag T Cell Responses in Clade B HIV-Infected Subjects



PCT/US01/28861

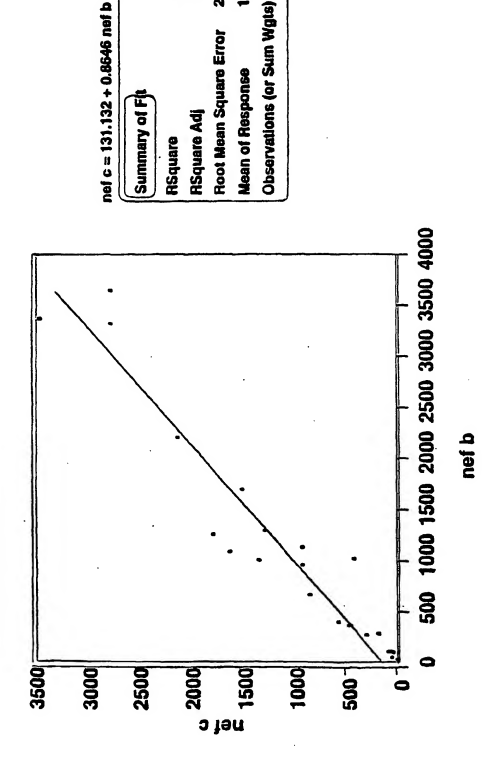
FIGURE 25

23

0.91685

289.7718 1096.435

Comparison of Clade B vs. Clade C Anti-nef T Cell Responses in Clade B HIV-Infected Subjects

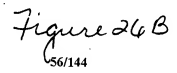


MRKAd5pol MER1062 (MRKAd5 Pre-Adenoviral Vector Containing the IA opt pol Coding Region)

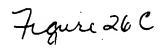
1	CATCATCAAT			GAAGCCAATA CTTCGGTTAT	
51	GGGGTGGAGT				
				ACCCTTGCCC	
101	TAGTAGTGTG ATCATCACAC			GTGTGGCGGA CACACCGCCT	
151				GTGTGCGCCG CACACGCGGC	
201				GATGTTGTAG CTACAACATC	
251	CGTAACCGAG GCATTGGCTC			GGGAAAACTG CCCTTTTGAC	
301				TAGCGCGTAA ATCGCGCATT	
351				AGACTCGCCC TCTGAGCGGG	
401				TTGGCGTTTT AACCGCAAAA	
451				CATATCATAA GTATAGTATT	
501				TGTTGACATT ACAACTGTAA	
551					AGCCCATATA TCGGGTATAT
601					TGGCTGACCG ACCGACTGGC
651	CCCAACGACC GGGTTGCTGG	CCCGCCCATT GGGCGGGTAA	GACGTCAATA CTGCAGTTAT	ATGACGTATG TACTGCATAC	TTCCCATAGT AAGGGTATCA
					TATTTACGGT ATAAATGCCA
751					AAGTACGCCC TTCATGCGGG
801					ATGCCCAGTA TACGGGTCAT
851					GTATTAGTCA CATAATCAGT

7 i jure 26A

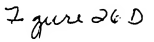
•					1017
901	TCGCTATTAC	CATGGTGATG	CGGTTTTGGC	AGTACATCAA	TGGGCGTGGA
	AGCGATAATG	GTACCACTAC	GCCAAAACCG	TCATGTAGTT	ACCCGCACCT
951	TAGCGGTTTG	ACTCACGGGG	ATTTCCAAGT	CTCCACCCCA	TTGACGTCAA
	ATCGCCAAAC	TGAGTGCCCC	TAAAGGTTCA	GAGGTGGGGT	AACTGCAGTT
1001	TGGGAGTTTG	TTTTGGCACC	AAAATCAACG	GGACTTTCCA	AAATGTCGTA
			_	CCTGAAAGGT	
1051				GTAGGCGTGT CATCCGCACA	
1101	GTCTATATAA	GCAGAGCTCG	TTTACTCAAC	CGTCAGATCG	CCTGGAGACG
				GCAGTCTAGC	
1151	CCATCCACGC	TGTTTTGACC	TCCATAGAAG	ACACCGGGAC	CGATCCAGCC
				TGTGGCCCTG	
1201	TCCGCGGCCG	GGAACGGTGC	ATTGGAACGC	GGATTCCCCG	TGCCAAGAGT
	AGGCGCCGGC	CCTTGCCACG	TAACCTTGCG	CCTAAGGGGC	ACGGTTCTCA
1251	GAGATCTACC	ATGGCCCCCA	TCTCCCCCAT	TGAGACTGTG	CCTGTGAAGC
	CTCTAGATGG	TACCGGGGGT	AGAGGGGGTA	ACTCTGACAC	GGACACTTCG
1301	TGAAGCCTGG	CATGGATGGC	CCCAAGGTGA	AGCAGTGGCC	CCTGACTGAG
	ACTTCGGACC	GTACCTACCG	GGGTTCCACT	TCGTCACCGG	GGACTGACTC
1351	GAGAAGATCA	AGGCCCTGGT	GGAAATCTGC	ACTGAGATGG	AGAAGGAGGG
	CTCTTCTAGT	TCCGGGACCA	CCTTTAGACG	TGACTCTACC	TCTTCCTCCC
1401	CAAAATCTCC	AAGATTGGCC	CCGAGAACCC	CTACAACACC	CCTGTGTTTG
				GATGTTGTGG	
1451				GGAAGCTGGT	
				CCTTCGACCA	
1501				GAGGTGCAGC	
				CTCCACGTCG	
1551	CCACCCGCT				
				ACACTGACAC	
1601	GGGATGCCTA CCCTACGGAT	• •		AGGACTTCAG TCCTGAAGTC	
1651				ACCCCTGGCA TGGGGACCGT	
1701	•••••	-		CTCCCCTGCC GAGGGGACGG	
1751	CCTCCATGAC GGAGGTACTG			GGAAGCAGAA CCTTCGTCTT	
1801	GTGATCTACC CACTAGATGG			GTGGGCTCTG CACCCGAGAC	



1901	GGGGCCTGAC CCCCGGACTG	CACCCCTGAC GTGGGGACTG			
1951		ATGAGCTGCA TACTCGACGT			
2001		AAGGACTCCT TTCCTGAGGA			
2051	GCAAGCTGAA CGTTCGACTT	CTGGGCCTCC GACCCGGAGG			
2101		TGCTGAGGGG ACGACTCCCC			
2151	GACTGAGGAG CTGACTCCTC	GCTGAGCTGG CGACTCGACC	AGCTGGCTGA TCGACCGACT	GAACAGGGAG CTTGTCCCTC	ATCCTGAAGG TAGGACTTCC
2201		TGGGGTGTAC ACCCCACATG			
2251	TAGGTCTTCG	AGGGCCAGGG TCCCGGTCCC	GGTCACCTGG .	ATGGTTTAGA	TGGTCCTCGG
2301	GAAGTTCTTG	CTGAAGACTG GACTTCTGAC	CGTTCATACG	GTCCTACTCC	CCCCGGGTGT
2351	GGTTACTACA	GAAGCAGCTG CTTCGTCGAC	TGACTCCGAC	ACGTCTTCTA	GTGGTGACTC
2401	AGGTAACACT	TCTGGGGCAA AGACCCCGTT	CTGGGGGTTC	AAGTTCGACG	GGTAGGTCTT
2451	CCTCTGGACC	GAGACCTGGT CTCTGGACCA	CCTGACTCAT	GACCGTCCGG	TGGACCTAGG
2501	GACTCACCCT	GTTTGTGAAC CAAACACTTG	TGGGGGGGG	ACCACTTCGA	CACCATGGTC
		TCGGGTAACA	CCCCGACTC	TGGAAGATAC	ACCGACCCCG
		CTCTGGTTCG	ACCCGTTCCG	ACCGATACAC	TGGTTGTCCC
		CCACCACTGG	GACTGACTGT	GGTGGTTGGT	CTTCTGACGG
2701	CTCCAGGCCA GAGGTCCGGT	TCTACCTGGC AGATGGACCG	CCTCCAGGAC GGAGGTCCTG	TCTGGCCTGG AGACCGGACC	AGGTGAACAT TCCACTTGTA
2751	TGTGACTGCC ACACTGACGG	TCCCAGTATG AGGGTCATAC	CCCTGGGCAT GGGACCCGTA	CATCCAGGCC GTAGGTCCGG	CAGCCTGATC GTCGGACTAG



2851			CACAAGGGCA GTGTTCCCGT	
2901			CATCAGGAAG GTAGTCCTTC	
2951			ATGAGAAGTA TACTCTTCAT	
3001			CCCCTGTGG GGGGGACACC	
3051			GAAGGGGGAG CTTCCCCCTC	
3101			AGCTGGCCTG TCGACCGGAC	
3151			GTGGCCTCCG CACCGGAGGC	
3201			GGAGACTGCC CCTCTGACGG	
3251			CCATCCACAC GGTAGGTGTG	
3301			GCCTGCTGGT CGGACGACCA	
3351			CCAGTCCCAG GGTCAGGGTC	
3401	CCTCCATGAA GGAGGTACTT		TTGGGCAGGT AACCCGTCCA	
3451			GCTGTGTTCA CGACACAAGT	
3501	CAAGAGGAAG GTTCTCCTTC		CGCTGGGGAG GCGACCCCTC	
3551	ACATCATTGC TGTAGTAACG		AGCTCCAGAA TCGAGGTCTT	
3601	AAGATCCAGA TTCTAGGTCT		GACTCCAGGA CTGAGGTCCT	
3651	GAAGGGCCCT CTTCCCGGGA		GGAGGGGGCT CCTCCCCGA	
3701	AGGACAACTC TCCTGTTGAG		GGAGGAAGGC CCTCCTTCCG	



3801		TAAAGCCCGG ATTTCGGGCC			
3851		TTGCCCCTCC AACGGGGAGG			
3901.	ACTCCCACTG TGAGGGTGAC	TCCTTTCCTA AGGAAAGGAT			
3951		CATTCTATTC GTAAGATAAG			
4001	CCCTCCTAAC	GGAAGACAAT CCTTCTGTTA	TCGTCCGTAC	GACCCCTACG	CCACCCGAGA
4051	TACCGGCTAG	GGCGCGCCGT CCGCGCGCA	TGACTTTACA	CACCCGCACC	GAATTCCCAC
4101	CCTTTCTTAT	TATAAGGTGG ATATTCCACC	CCCAGAATAC	ATCAAAACAT	AGACAAAACG
4151	TCGTCGGCGG	GCCGCCATGA CGGCGGTACT	CGTGGTTGAG	CAAACTACCT	TCGTAACACT
4201	CGAGTATAAA	GACAACGCGC CTGTTGCGCG	TACGGGGGTA	CCCGGCCCCA	CGCAGTCTTA
4251	CACTACCCGA	CCAGCATTGA GGTCGTAACT	ACCAGCGGGG	CAGGACGGGC	GTTTGAGATG
4301	ATGGAACTGG	TACGAGACCG ATGCTCTGGC	ACAGACCTTG	CGGCAACCTC	TGACGTCGGA
4351	GGCGGCGGCG	TTCAGCCGCT AAGTCGGCGA	CGTCGGTGGC	GGGCGCCCTA	ACACTGACTG
4401	AAACGAAAGG		ACGTTTGTCA	CGTCGAAGGG	CAAGTAGGCG
	GGCGCTACTG	TTCAACTGCC	GAGAAAACCG	TGTTAACCTA	TCTTTGACCC AGAAACTGGG
	CCCTTGAATT	ACAGCAAAGA	GTCGTCGACA	ACCTAGACGC	CCAGCAGGTT GGTCGTCCAA
	AGACGGGACT	TCCGAAGGAG	GGGAGGGTTA	CGCCAAATTT	ACATAAATAA TGTATTTATT
	TTTTGGTCTG	AGACAAACCT	AAACCTAGTT	CGTTCACAGA	TGCTGTCTTT ACGACAGAAA
4651					GTCTCGGTCG CAGAGCCAGC

Figure 26E 59/144

4751				GGGGTGGAGG CCCCACCTCC	
4801	GCAGAGCTTC CGTCTCGAAG	ATGCTGCGGG TACGACGCCC	GTGGTGTTGT CACCACAACA	AGATGATCCA TCTACTAGGT	GTCGTAGCAG CAGCATCGTC
4851				TTCAGTAGCA AAGTCATCGT	
4901				AAAGCGGTTA TTTCGCCAAT	
4951	CCACGTATGC	ACCCCTATAC	TCTACGTAGA	TGGACTGTAT ACCTGACATA	AAAATCCAAC
5001	CGATACAAGG	GTCGGTATAG	GGAGGCCCCT	TTCATGTTGT AAGTACAACA	CGTCTTGGTG
5051	GTCGTGTCAC	ATAGGCCACG	TGAACCCTTT	TTTGTCATGT AAACAGTACA	TCGAATCTTC
5101	CTTTACGCAC	CTTCTTGAAC	CTCTGCGGGA	TGTGACCTCC ACACTGGAGG	TTCTAAAAGG
5151	TACGTAAGCA	GGTATTACTA	CCGTTACCCG	CCACGGGCGG GGTGCCCGCC	GCCGGACCCG
5201	СТТСТАТААА	GACCCTAGTG	ATTGCAGTAT	GTTGTGTTCC CAACACAAGG	TCCTACTCTA
5251	GCAGTATCCG	GTAAAAATGT	TTCGCGCCCG	GGAGGGTGCC CCTCCCACGG	TCTGACGCCA
5301	TATTACCAAG	GTAGGCCGGG	TCCCCGCATC	TTACCCTCAC AATGGGAGTG	TCTAAACGTA
5351	AAGGGTGCGA	AACTCAAGTC	TACCCCCTA	CATGTCTACC GTACAGATGG	ACGCCCCGCT
	ACTTCTTTTG	CCAAAGGCCC	CATCCCCTCT	AGTCGACCCT	AGAAAGCAGG TCTTTCGTCC
	AAGGACTCGT	CGACGCTGAA	TGGCGTCGGC	CACCCGGGCA	AAATCACACC TTTAGTGTGG
	ATAATGGCCG	ACGTTGACCA	TCAATTCTCT	CGACGTCGAC	CCGTCATCCC GGCAGTAGGG
	ACTCGTCCCC	CCGGTGAAGC	AATTCGTACA	GGGACTGAGC	CATGTTTTCC GTACAAAAGG
5601					GCAGTTCTTG CGTCAAGAAC

Figure 26 F

5701		TTGACCAAGC AACTGGTTCG			
5751		CATCTCGATC GTAGAGCTAG			
5801		TGTACGGCAG ACATGCCGTC			
5851		CACGGGCGCA GTGCCCGCGT			
5901		CGCTCCGGGC GCGAGGCCCG			
5951	GTCCTGCTGG CAGGACGACC	TGCTGAAGCG ACGACTTCGC	CTGCCGGTCT GACGGCCAGA	TCGCCCTGCG AGCGGGACGC	CGTCGGCCAG GCAGCCGGTC
6001		ACCATGGTGT TGGTACCACA			
6051	TGGCGCGCAG ACCGCGCGTC	CTTGCCCTTG GAACGGGAAC	GAGGAGGCGC CTCCTCCGCG	CGCACGAGGG GCGTGCTCCC	GCAGTGCAGA CGTCACGTCT
6101	CTTTTGAGGG GAAAACTCCC	CGTAGAGCTT GCATCTCGAA	GGGCGCGAGA CCCGCGCTCT	AATACCGATT TTATGGCTAA	CCGGGGAGTA GGCCCCTCAT
6151		CCGCAGGCCC GGCGTCCGGG			
6201		CCGTTCGGGG GGCAAGCCCC			
6251		TACCTCTGGT ATGGAGACCA			
6301		TCCGTGTCCC AGGCACAGGG			
6351					CTCTGAGACA GAGACTCTGT
6401	AAGGCTCGCG TTCCGAGCGC	TCCAGGCCAG AGGTCCGGTC	CACGAAGGAG GTGCTTCCTC	GCTAAGTGGG CGATTCACCC	AGGGGTAGCG TCCCCATCGC
6451					AGACACATGT TCTGTGTACA
6501					GTAGGCCACG CATCCGGTGC
6551	TGACCGGGTG ACTGGCCCAC	TTCCTGAAGG AAGGACTTCC	GGGGCTATAA CCCCGATATT	AAGGGGGTGG TTCCCCCACC	GGGCGCGTTC CCCGCGCAAG

Figure 266

6651		CTGAAAAGCG GACTTTTCGC			
6701		AGGAGGATTT TCCTCCTAAA			
6751		GCATCCATCT CGTAGGTAGA			
6801		AAACGACCCG TTTGCTGGGC			
6851		TTTGGTTTTT AAACCAAAAA			
6901	CAAATCGACG	ACGTATTCGC TGCATAAGCG	CGCGTTGCGT	GGCGGTAAGC	CCTTTCTGCC
6951	ACCACGCGAG	GTCGGGCACC CAGCCCGTGG	TCCACGTGCG	CGGTTGGCGC	CAACACGTCC
7001	CACTGTTCCA	CAACGCTGGT GTTGCGACCA	CCGATGGAGA	GGCGCATCCG	CGAGCAACCA
7051	GGTCGTCTCC	CGGCCGCCCT GCCGGCGGGA	ACGCGCTCGT	CTTACCGCCA	TCCCCCAGAT
7101	CGACGCAGAG	GTCCGGGGGG CAGGCCCCCC	AGACGCAGGT	GCCATTTCTG	GGGCCCGTCG
7151	TCCGCGCGCA	CGAAGTAGTC GCTTCATCAG	ATAGAACGTA	GGAACGTTCA	GATCGCGGAC
7201	GACGGTACGC	CGGGCGGCAA GCCCGCCGTT	CGCGCGCGAG	CATACCCAAC	TCACCCCCTG
7251	GGGTACCGTA	GGGGTGGGTG CCCCACCCAC	TCGCGCCTCC	GCATGTACGG	CGTTTACAGC
	ATTTGCATCT	CCCCGAGAGA	CTCATAAGGT	TCTATACATC	GGTAGCATCT CCATCGTAGA
		TACGACCGCG	CGTGCATTAG	CATATCAAGC	ACGCTCCCTC
		CCCTGGCTCC	AACGATGCCC	GCCCGACGAG	ACGAGCCTTC
		ACTTCTACCG	TACACTCAAC	CTACTATACC	AACCTGCGAC
7501	GAAGACGTTG CTTCTGCAAC	AAGCTGGCGT TTCGACCGCA	CTGTGAGACC GACACTCTGG	TACCGCGTCA ATGGCGCAGT	CGCACGAAGG GCGTGCTTCC

Figure 26 H

7601				ATGATGTCAT TACTACAGTA	
7651				GACAAACTCT CTGTTTGAGA	
7701				CCTCCGAACG GGAGGCTTGC	
7751				GCGCAGCATC CGCGTCGTAG	
7801				GAGCGAGGTG CTCGCTCCAC	
7851				ACTGGTATTT TGACCATAAA	
7901				AAGTCCGTGC TTCAGGCACG	
7951				GTTGAAGAGT CAACTTCTCA	
8001				AGGGTCCCGG TCCCAGGGCC	
8051				ATCTCGTCAA TAGAGCAGTT	
8101				GCGCGGGATG CGCGCCCTAC	
8151				GCTCTTCAGG CGAGAAGTCC	
8201	CCGTGCTCTG GGCACGAGAC	AAAGGGCCCA TTTCCCGGGT	GTCTGCAAGA CAGACGTTCT	TGAGGGTTGG ACTCCCAACC	AAGCGACGAA TTCGCTGCTT
8251	TGAGCTCCAC ACTCGAGGTG			TTGCAGGTGG AACGTCCACC	
8301	TCCTAAACTG AGGATTTGAC			CTGGGGTGAT GACCCCACTA	
8351	GTAAGCGGGT CATTCGCCCA			CCAAGGTTCG GGTTCCAAGC	
	TCGCGCGGCA AGCGCGCCGT				
8451	TGAAGGGCAC ACTTCCCGTG			CCATCCAAGT GGTAGGTTCA	

Figure 26I

8551				GGAGTGGCTA CCTCACCGAT	
8601	GAAAGTAGAA CTTTCATCTT	GTCCCTGCGA CAGGGACGCT	CGGGCCGAAC GCCCGGCTTG	ACTCGTGCTG TGAGCACGAC	GCTTTTGTAA CGAAAACATT
8651	AAACGTGCGC TTTGCACGCG	AGTACTGGCA TCATGACCGT	GCGGTGCACG CGCCACGTGC	GGCTGTACAT CCGACATGTA	CCTGCACGAG GGACGTGCTC
8701				GAGTGGGAAT CTCACCCTTA	
8751			TGGTCTTCTA ACCAGAAGAT	CTTCGGCTGC GAAGCCGACG	TTGTCCTTGA AACAGGAACT
8801				GATCGGACCA CTAGCCTGGT	
8851			CCGCGCGCGC	CGGTCGGAGC GCCAGCCTCG	TTGATGACAA AACTACTGTT
8901				GGAGCTCCCG CCTCGAGGGC	
8951				CATAGACGGG GTATCTGCCC	
9001		AGGTGATACC TCCACTATGG		GGGCTGGTTG CCCGACCAAC	
9051			CATCCCCGCG GTAGGGGCGC	GCGCGACTAC CGCGCTGATG	GGTACCGCGC CCATGGCGCG
9101			GGTGTCCTTG CCACAGGAAC	GATGATGCAT CTACTACGTA	CTAAAAGCGG GATTTTCGCC
9151	TGACGCGGGC ACTGCGCCCG	GAGCCCCCGG CTCGGGGGCC	AGGTAGGGGG TCCATCCCCC	GGCTCCGGAC CCGAGGCCTG	CCGCCGGGAG GGCGGCCCTC
9201	AGGGGGCAGG TCCCCCGTCC	GGCACGTCGG CCGTGCAGCC	CCCCCCCCCC	GGGCAGGAGC CCCGTCCTCG	TGGTGCTGCG ACCACGACGC
9251	CGCGTAGGTT GCGCATCCAA	GCTGGCGAAC CGACCGCTTG	GCGACGACGC CGCTGCTGCG	GGCGGTTGAT CCGCCAACTA	CTCCTGAATC GAGGACTTAG
9301	TGGCGCCTCT ACCGCGGAGA	GCGTGAAGAC CGCACTTCTG	GACGGGCCCG CTGCCCGGGC	GTGAGCTTGA CACTCGAACT	ACCTGAAAGA TGGACTTTCT
9351	GAGTTCGACA CTCAAGCTGT	GAATCAATTT CTTAGTTAAA	CGGTGTCGTT GCCACAGCAA	CTCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC	TGGCGCAAAA ACCGCGTTTT
9401	TCTCCTGCAC AGAGGACGTG	GTCTCCTGAG CAGAGGACTC	TTGTCTTGAT AACAGAACTA	AGGCGATCTC TCCGCTAGAG	GGCCATGAAC CCGGTACTTG

Figure 26 J

9501				GAGCTGCGAG CTCGACGCTC	
9551				CCACGCCCCC GGTGCGGGGG	
9601				AGCTCCACGT TCGAGGTGCA	
9651				GTAGTTGAGG CATCAACTCC	
9701				AGCGTCGCAA TCGCAGCGTT	
9751		GGTTCCGGAG	TTCCGCGAGG	TACCGGAGCA	TCTTCAGGTG
9801				CGACACGGTT GCTGTGCCAA	
9851				CGCGCACCTC GCGCGTGGAG	
9901				TCCTCTTCCA AGGAGAAGGT	
9951	GGGAAGAAGA	AGAAGACCGC	CGCCACCCCC	AGGGGGGACA TCCCCCTGT	GCCGCCGCTG
10001				GCTCGATCAT CGAGCTAGTA	
10051	GCTGCCGCGT	ACCAGAGCCA	CTGCCGCGCC	CCGTTCTCGC GGCAAGAGCG	CCCCGCGTC
10101				ATGGGTTGGC TACCCAACCG	
10151	CATGCGGCAG GTACGCCGTC				TTGTTGTGTA AACAACACAT
		GCGGCTCCCT	GGACTCGCTC	AGGCGTAGCT	GGCCTAGCCT
10251	AAACCTCTCG TTTGGAGAGC			ACAGTCGCAA TGTCAGCGTT	
10301	GCACCGTGGC CGTGGCACCG	GGGCGGCAGC CCCGCCGTCG	GGGCGGCGT	CGGGGTTGTT	TCTGGCGGAG AGACCGCCTC
10351	GTGCTGCTGA CACGACGACT				GGCGGATGGT CCGCCTACCA

Figure 26 K

10451	CGGCCATGCC GCCGGTACGG			GGCGCAGGTC CCGCGTCCAG	
10501				TCTTCTCCTT AGAAGAGGAA	
10551				GGCGGAGTTT CCGCCTCAAA	
10601				CGAAGCCCCT GCTTCGGGGA	
10651				GCTAATATGG CGATTATACC	
10701	CTGCGTGAGG GACGCACTCC			GTCCACAAAG CAGGTGTTTC	
10751				CCATAACGGA GGTATTGCCT	
10801				TACCTGAGAC ATGGACTCTG	
10851				CCGCACCAGG GGCGTGGTCC	
10901				AGAGGGGCCA TCTCCCCGGT	
10951				ATAAGGCGAT TATTCCGCTA	
11001	CTACATGGAC	CTGTAGGTCC	ACTACGGCCG	GGCGGTGGTG CCGCCACCAC	CTCCGCGCGC
11051				GCAGCGGCAA CGTCGCCGTT	
	TACCAGCCCT	GCGAGACCGG	CCAGTCCGCG	CGCGTTAGCA	TGACGCTCTA ACTGCGAGAT
		TTCCTCTCGG	ACATTCGCCC	GTGAGAAGGC	ACCAGACCAC
	CTATTTAAGC	GTTCCCATAG	TACCGCCTGC	TGGCCCCAAG	GAGCCCCGTA CTCGGGGCAT
		GCGGCACTAG	GTACGCCAAT	GGCGGGCGCA	CAGCTTGGGT
11301					TTCCTTCCAG AAGGAAGGTC

Figure 26L

11401		GCTGGAAAGC CGACCTTTCG		
11451		TTTTCCAAGG AAAAGGTTCC		
11501		GGACTGCGGC CCTGACGCCG		
11551		CAAATTCCTC GTTTAAGGAG		
11601		CATCCGGTGC GTAGGCCACG		
11651		AGAGCAGCGG TCTCGTCGCC		
11701		GAGGGGCGAC CTCCCCGCTG		
11751		GCGCGCGCC		
11801		GGCGCGGCTA CCGCGCCGAT		
11851		AGCGTGATAC TCGCACTATG		
11901		CGCGAGGGAG GCGCTCCCTC		
11951		GCGCGAGCTG CGCGCTCGAC		
12001		ACTTTGAGCC TGAAACTCGG		
12051	CGCACACGTG GCGTGTGCAC	922992999 922992999		
12101	ACCAGGAGAT TGGTCCTCTA	TAACTTTCAA ATTGAAAGTT		
12151	GTGGCGCGCG CACCGCGCGC	AGGAGGTGGC TCCTCCACCG		
12201	AAGCGCGCTG TTCGCGCGAC	GAGCAAAACC CTCGTTTTGG		
12251	TCCTTATAGT AGGAATATCA	GCAGCACAGC CGTCGTGTCG		



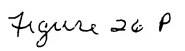
TACCECCEC TATCACCACE TECTEGGTE GAACTGGAC CGACTGTTCC 12401 TGGCCGCGTA CAACTATTCC ATGCTTAGCC TGGGCAAGTT TTACGCCGGC 12451 AAGATATACC ATACCCCTTA CGTTCCCATA GACAAGGGGG TAAAGATCGA 12501 GGGGTTCTAC ATGCCCATG CGCGCACGACGT CTGCTCACCACACACCCCCCAAGATG TACGCATGC CCCCAAGATG TACGCGTACC CCCCACCACA AGGCCCTGCACCC CCACCCACCACA ACCCCGCCACCACA ACCCCCCCC	12351				CTTGAGCCTG	
ACCGGCGGTA GTTGATAAGG TACGAATCGG ACCCGTTCAA AATGCGGCG 12451 AAGATATACC ATACCCCTTA CGTTCCCATA GACAAGGAGG TAAAGATCGA TTCTATATGG TATGGGGAAT GCAAGGGTAT CTGTTCCTCC ATTTCTAGCT 12501 GGGGTTCTAC ATGCGCATGG CGCTGAAGGT GCTTACCTTG AGCGACGAC CCCCAAGATG TACGCGTACC GCGACTCCAC AGGCCGTAAG 12551 TGGGCGTTTA TCGCAACGAG CGCATCCACA AGGCCGTGAG CGTGAGCCG ACCCGCAAAT AGCGTTGCTC GCGTAGGTT TCCGGCACTC GCACTCGGC 12601 CGGCGGGAGC TCAGCGACCG CGAGCTGATC CACAGCCTGC AAAGGGCCCT GCCGCGCTCG AGTCGCTGCC GCTCAGCACA GTTCCGCACCT CCGACCGGC 12701 GCGCTGACCG GGCAGCGGC ATAGAAGAGGC CGAGTCCTAC TTTCACGCGC 12701 GCGCTGACCT GCGCTGGCCG TATCTCTCG GCTCAGGATG AAACTGCGCC 12771 GCGCTGACCT GCGCTGGCCG TATCTCTCG GCTCAGGATG AAACTGCGCC 12751 GCCGGACCTG GGCTGGCCG CCAAGCCGAC GCGCCCTGGA CCGCCCCCCC CCGCGACTGGA CCGGACCCGC GGTTCGGCTG CGCGGGACCT CCGTCGACCC 12751 GCCGGACCTG GGCTGGCCG CCAAGCCGAC GCGCCGTGGA AACTGCGCC 12801 CGTGGAGGAA TATGACGAGG ACGATCCTAC TTTGACGCGG CCGCCTCTT ATACTGCTCC TGCTAGCTTAC TCAGGCCGC 12851 ACTAAGCGGT GATGTTTCT ATCAGAGGA CCGAGCCAGAG GACCCGCCCCC 12901 CGGTGGGGC GATGTTTCT ATCAGATGAT CGAGCCAGAG GACCGCGCACT TGATTCGCCA CTACAAAGAC TAGCTACTA GCTCGGTCT TGCCTGGCCT 12901 CGGTGGGGC GCGCTGCAG AGCCAGCCGT CCGGCCGAATT GACCGCCCC 12901 CGGTGGGGC GCGCTGCAG AGCCAGCCGT CCGGCCTTAA CTCCACGGAC GCCACGCCCG CCGCGACGTC TCGGTCGCA GCCCGGAATT GACGCCCGC 12901 CGGTGGGGC AGCTCAGAA CCGCACCACA GCCCGCAATC CCTGACCGCG TCCAGTACCT GGCTGGCAC GCCCGGAATT GACGCCCGC CCGCACGCCG CCGCGACGTC TCGGTCGCC CCGCCAATC CCTGACCGCG TCCAGTACCT GGCCTGAAA GCGCAATCC CTGACCGCGG TCCAGTACCT GGCCTGGAC GCCCGAATC CCGCCAACCCCG CCGCGCCG CCGCACGCCA CCGCCAACAC 13001 TGACGCGTTC CGGCCGCC GCAAACCCCA CGCACGAAA AACGGCCGGC TTCAGCACCGC TCCAGCCCG CCGCACGCCA CCGCCAACAC 13001 TGACGCGTTC CGGCCGC GCGCACCCC CCGCACACCC CCGCACACCA AACCCCGC GCCACCGCC CCGCCCCC CCGCACCCC CCGCACACCA CCGCCCACACCA CCGCCCACCAC CCGCCACCAC CCGCCACCAC CCGCCCCCCCC						
12451 AAGATATACC ATACCCCTTA CGTTCCCATA GACAAGGAGG TAAAGATCGA TTCTATATAGG TATGGGAAT GCAAGGGTAT CTGTTCCTCC ATTTCTAGCT 12501 GGGGTTCTAC ATGCGCATGG CGCTGAAGGT CCTTACCTTG AGCGACGACC CCCCAAGATG TACGCCATACC GCGACTTCCA CGAATGGAAC TCGCTGCTGG 12551 TGGGCGTTTA TCGCAACGAG CGCATCCACA AGGCCGTAGA CGTGAGCCGG ACCCGCAAAT AGCGTTGCTC GCGATGGTT TCCGGGACTC GCACTCGGCC 12601 CGGCGCGAGC TCAGCGACCG CGAGCTGAT CACAGCCTGC AAAGGCCCT AGCCGGCACCGC CCGCCCCCC AGCTCGCC CCGCCCCC ATTCCCCGGA 12651 GGCTGGCACG GGCAGCGGCG ATAGAGAGGC CGAGTCCTAC TTTGACGCGG CCGACCGTGC CCGTCGCCG TATCTCTCCG GCTCAGGATG AAACTGCGCC 12701 GCGCTGACCT GCGCTGGCC CCAAGCCGAC GCCCCTGGA GGCACCTGGC CCGCGACTGGA CGCGACCGCG GGTTCGGCT CCGTCGGACCTC CCGCGACTGGA CGCGACCGCG GGTTCGGCT CCGTCGGAC CCGTCGACCCC 12751 GCCGGACCTG GCGTGGCGC CCAAGCCGAC GCCCCTGGA GGCAGCTGGC CCGCCTGGAC CCGACCGCG GGTTCGGCT CCGTCGGAC CCGTCGACCCC 12801 CGTGGAGGAA TATGACGAGG ACGATCAGTA CGCCCCCCC 12801 CGTGGAGGAA TATGACGAGG ACGATCAGTA CGCCCCGCCC	12401					
TTCTATATGG TATGGGGAAT GCAAGGGTAT CTGTTCCTC ATTTCTAGCT 12501 GGGGTTCTAC ATGCGCATGG CGCTGAAGGT GCTTACCTTG AGCGACGACC CCCCAAGATG TACGCGTACC GCGATCCACA CGAATGGAAC TCGCTGCTGG ACCCGCAAAT ACCGTTGCTC GCGTAGGTGT TCCGGCACCTC GCACCGCCAAAT ACCGTTGCTC GCGTAGGTGT TCCGGCACCTC GCACCTGGC ACCCGCAAAT ACCGTTGCTC GCGTAGGTGT TCCGGCACCTG GCCGCGCCTC AATGCGACCG CGAGCTGATG CACAGCCTGC AAAGGGCCCT GCGCGCCTCG AATCGCTGCC CGTCGACCTAC GTGTCGGACCT GCCGCCCCCC AATCCCTCAC ATTCCCGGAACTAC GTGTCGGACC CCGACCGTGC CCGTCGCCC TATCTCTCCG GCTCAGGATG AAACTGCGCC CCGACCGTGC CCGTCGCCC TATCTCTCCG GCTCAGGATG AAACTGCGCC CCGACCGTGC CCGTCGCCC TATCTCTCCG GCTCAGGATG AAACTGCGCC CCGCGACCGTGC CCGTCGCCGC CCAAGCCGAC GCGCCCTGGA GGCACCGCG GGTTCGGCT CCGTCGACCT CCGTCGACCT CCGTCGACCT CCGGCACCTGA CCGCGCCCCC CCGACCGCCA CCGGCGCCCC CCGCGCACCGCC CCGACCGCCA CCGGCGCACCGC GCGCGGACCGT TGCAGCCCCC CCGACCGCCA CCGGCCACGCCA CCGGCGCACCGCT TGCAGCCGC CCGACCGCCA CCGACCGCCA CCGGCGCACCGT TGCAGCCGCC CCGACCGCCA CCGACCGCCA CCGGCGCACCGCT TGCAGCCGCC ACGGCCACCGCC ACGACCACCCC CCGACCGCCA CCGACCGCAC CCGACCGCCA CCGCCCACCGCC CCCCCACCGCC CCCCCACCACCC CCCCCCACCCCC CCCCCACCACC		ACCGGCGGTA	GTTGATAAGG	TACGAATCGG	ACCCGTTCAA	AATGCGGGCG
12501 GGGGTTCTAC ATGCGCATGG CGCGCTGAAGGT GCTTACCTTG AGCGACGAC CCCCAAGATG TACGCGTACC GCGACTTCCA CGAATGGAAC TCGCTGCGGCCCCCCCCAAAT AGCGTTGCTC GCGATCCACA AGGCCGTAGA CGCACTCGGCCCCCCCCCC	12451					
CCCCAAGATG TACGCGTACC GCGACTTCCA CGAATGGAAC TCGCTGCTGG 12551 TGGGCGTTTA TCGCAACGAG CGCATCCACA AGGCCGTGAG CGTGAGCCGG ACCCGCAAAT AGCGTTGCTC GCGTAGGTGT TCCGGCACTC GCACTCGGCC 12601 CGGCGCGAGC TCAGCGACCG CGAGCTGATG TCCGGCACTC GCACTCGGCC 12651 GGCTGGCACG GGCAGCGGCG ATAGAGAGGC CGAGTCCTAC TTTCCCGGGA 12701 GCGCTGACCT CCGCTGCCGC TATCTCTCCG GCTCAGGATG AAACTGCGCC 12701 GCGCTGACCT GCGCTGGCCC CCAAGCCGAC GCGCCCTGGA GGCACTGGC 12752 CCGGACCTG GCCTGCCGC GGTTCGGCC CGCGGACCCT CCGTCGACCC 12751 CCGGGACCTG GCCTGCCGC GGTTCGGCC GCGCGGACCCT TGCAGCCC 12801 CGTGGAGGAA TATGACGAGG ACGATCACAT GCCGGACCGC CCGGCCCCGCACCCCC ACGACCCACAT TGCAGCCGCC 12851 ACTAAGCGGT GATGTTTCTG ATCAGATGAT GCAAGACGCA ACGGACCGGC 12901 CGGTGCGGGC GGCGCTGCAA ACCAGCCGT CCGCCCTCAA 12901 CGGTCGGGCC GCGCGCACGTC TCGGTCGCC CCGCCACCCC 12951 GACTGGCGC GCGCGCACCGC AGCCACCGT TGCCCGGCC 12951 GACTGGCGC AGGTCATGA AGCCAGCCGT CCGCCCTAAAGACCCAC CCGCACCCCC CCGCACCGC CCGCACCCC CCCCCACCCCC CCCCCACCCC CCCCCACCCC CCCCCC						
CCCCAAGATG TACGCGTACC GCGACTTCCA CGAATGGAAC TCGCTGCTGG 12551 TGGGCGTTTA TCGCAACGAG CGCATCCACA AGGCCGTGAG CGTGAGCCGG ACCCGCAAAT AGCGTTGCTC GCGTAGGTGT TCCGGCACTC GCACTCGGCC 12601 CGGCGCGAGC TCAGCGACCG CGAGCTGATG TCCGGCACTC GCACTCGGCC 12651 GGCTGGCACG GGCAGCGGCG ATAGAGAGGC CGAGTCCTAC TTTCCCGGGA 12701 GCGCTGACCT CCGCTGCCGC TATCTCTCCG GCTCAGGATG AAACTGCGCC 12701 GCGCTGACCT GCGCTGGCCC CCAAGCCGAC GCGCCCTGGA GGCACTGGC 12752 CCGGACCTG GCCTGCCGC GGTTCGGCC CGCGGACCCT CCGTCGACCC 12751 CCGGGACCTG GCCTGCCGC GGTTCGGCC GCGCGGACCCT TGCAGCCC 12801 CGTGGAGGAA TATGACGAGG ACGATCACAT GCCGGACCGC CCGGCCCCGCACCCCC ACGACCCACAT TGCAGCCGCC 12851 ACTAAGCGGT GATGTTTCTG ATCAGATGAT GCAAGACGCA ACGGACCGGC 12901 CGGTGCGGGC GGCGCTGCAA ACCAGCCGT CCGCCCTCAA 12901 CGGTCGGGCC GCGCGCACGTC TCGGTCGCC CCGCCACCCC 12951 GACTGGCGC GCGCGCACCGC AGCCACCGT TGCCCGGCC 12951 GACTGGCGC AGGTCATGA AGCCAGCCGT CCGCCCTAAAGACCCAC CCGCACCCCC CCGCACCGC CCGCACCCC CCCCCACCCCC CCCCCACCCC CCCCCACCCC CCCCCC	12501	GGGGTTCTAC	ATGCGCATGG	CGCTGAAGGT	GCTTACCTTG	AGCGACGACC
ACCCGCAAAT AGCGTTGCTC GCGTAGGTGT TCCGGCACTC GCACTCGGCC 12601 CGGCGCGAGC TCAGCGACCG CGAGCTGATC CACAGCCTGC AAAGGGCCCT GCCGCGCTCG AGTCGCTGGC GCTCGACTAC GTGTCGGACG TTTCCCGGGA 12651 GGCTGGCACG GGCAGCGGCG ATAGAGGAGGC CGAGTCCTAC TTTGACGCGG CCGACCGTGC CCGTCGCCGC TATCTCTCCG GCTCAGGATG AAACTGCGCC 12701 GCGCTGACCT GCGCTGGCGC CCAAGCCGAC GCGCCCTGGA GGCAGCTGGG CGCGACCTGG CGCGACCCGG GGTTCGGCT CGCGGGACCT CCGTCGACCC 12752 GCCGGACCTG GGCTGGCGGT GGCACCGCG CGCGCTGGCA ACGTCGGCC 12801 CGTGGAGGAA TATGACGAGG ACGATGAGTA CGAGCCAGAG GACGGCGGC 12851 ACTAAGCGGT GATGTTTCTG ATCAGATGAT CGACCCAGAG GACGGCAGCT TGATTCGCCA CTACAAAGAC TAGTCTACTA CGTTCTGCGT TGCCTGGGCC 12901 CGGTGCGGGC GGCGCTGCAG AGCCAGCCGT CCGGCCTTAA CTCCACGGAC GCCACCCCCC CCGCGACGTC TCGGTCGGC GCCCGGAATT GACTGGCCTG 12951 GACTGGCGCC AGGTCATGAA CCCGACCCGT CCGGCCTTAA CTCCACGGAC CTGACCGCGC CCGCGACGT TCGGTCGGCA GCCCGGAATCC CTGACCGCGG TCCAGTACCT GGCGTAGTA AGCGACCTGA GAGGTCCTG 13001 TGACGCGTC CGGCAGCAG CCGACGCAA CCGGCCTTAA GCCGCCTTAGG 13001 TGACGCGTTC CGGCAGCAGC CGCAGGCCAA CCGGCCTTAGGCC 13101 AACCGGTGGT CCCGCCGCC CGCAAACCCCA CGGCCGAAACC 13051 AACCGGTGGT CCCGGCGCC CGTTTGGGGT GCCTGCCG TTCGCCACCA GGCCGCGCC CGTTTGGGGT GCCGAGAAA GGTCCTGCCG 13101 ACCGTAAACG CCCGCCGCC CGTTTTGCCCCG ACGACGAA GGTCCTGCCG CCCACCCCC GCGACCGCC CGTTTTGGGGT GCCTGCCCACACCGC 13101 ACCGTAAACG CCCGCCGCC CGTTTTGCCCCG ACGACGAA GGTCCTGCCG TAGCACTTAC AACAGGCC ATCCGGCCCA AACACCCA ACGACCAA CCACGCCCA ACGAGCCAA CCCACGCCCGCC		CCCCAAGATG	TACGCGTACC	GCGACTTCCA	CGAATGGAAC	TCGCTGCTGG
ACCCGCAAAT AGCGTTGCTC GCGTAGGTGT TCCGGCACTC GCACTCGGCC 12601 CGGCGCGAGC TCAGCGACCG CGAGCTGATC CACAGCCTGC AAAGGGCCCT GCCGCGCTCG AGTCGCTGGC GCTCGACTAC GTGTCGGACG TTTCCCGGGA 12651 GGCTGGCACG GGCAGCGGCG ATAGAGGAGGC CGAGTCCTAC TTTGACGCGG CCGACCGTGC CCGTCGCCGC TATCTCTCCG GCTCAGGATG AAACTGCGCC 12701 GCGCTGACCT GCGCTGGCGC CCAAGCCGAC GCGCCCTGGA GGCAGCTGGG CGCGACCTGG CGCGACCCGG GGTTCGGCT CGCGGGACCT CCGTCGACCC 12752 GCCGGACCTG GGCTGGCGGT GGCACCGCG CGCGCTGGCA ACGTCGGCC 12801 CGTGGAGGAA TATGACGAGG ACGATGAGTA CGAGCCAGAG GACGGCGGC 12851 ACTAAGCGGT GATGTTTCTG ATCAGATGAT CGACCCAGAG GACGGCAGCT TGATTCGCCA CTACAAAGAC TAGTCTACTA CGTTCTGCGT TGCCTGGGCC 12901 CGGTGCGGGC GGCGCTGCAG AGCCAGCCGT CCGGCCTTAA CTCCACGGAC GCCACCCCCC CCGCGACGTC TCGGTCGGC GCCCGGAATT GACTGGCCTG 12951 GACTGGCGCC AGGTCATGAA CCCGACCCGT CCGGCCTTAA CTCCACGGAC CTGACCGCGC CCGCGACGT TCGGTCGGCA GCCCGGAATCC CTGACCGCGG TCCAGTACCT GGCGTAGTA AGCGACCTGA GAGGTCCTG 13001 TGACGCGTC CGGCAGCAG CCGACGCAA CCGGCCTTAA GCCGCCTTAGG 13001 TGACGCGTTC CGGCAGCAGC CGCAGGCCAA CCGGCCTTAGGCC 13101 AACCGGTGGT CCCGCCGCC CGCAAACCCCA CGGCCGAAACC 13051 AACCGGTGGT CCCGGCGCC CGTTTGGGGT GCCTGCCG TTCGCCACCA GGCCGCGCC CGTTTGGGGT GCCGAGAAA GGTCCTGCCG 13101 ACCGTAAACG CCCGCCGCC CGTTTTGCCCCG ACGACGAA GGTCCTGCCG CCCACCCCC GCGACCGCC CGTTTTGGGGT GCCTGCCCACACCGC 13101 ACCGTAAACG CCCGCCGCC CGTTTTGCCCCG ACGACGAA GGTCCTGCCG TAGCACTTAC AACAGGCC ATCCGGCCCA AACACCCA ACGACCAA CCACGCCCA ACGAGCCAA CCCACGCCCGCC	12551	TGGGCGTTTA	TCGCAACGAG	CGCATCCACA	AGGCCGTGAG	CGTGAGCCGG
12601 CGGCGGAGC TCAGCGACCG GCTCGACTAC GTGTCGGACG TTTCCCGGGA 12651 GGCTGGCACG GGCAGCGGCG ATAGAGAGGC CGAGTCCTAC TTTGACGCGG 12701 GCGCTGACCT GCGCTGGCCG CAAGCCGAC GCTCAGGATG AAACTGCGCC 12771 GCGGTGACCT GCGCTGGCCG CCAAGCCGAC GCGCCCTGGA GGCAGCTGGA CGCGACCGGG GGTTCGGCTG CGCGGACCCG GGTTCGGCCG CGCGGACCCG GGTCCGGACCCG GGTTCGGCGC CGCGGACCGG CCGGCCTGGA ACGTCGGCGC CGGCCTGGAC CCGGCCTGGAC CCGGCCTGGAC CCGGCCTGGAC CCGGCCTGGACCCC CGGCCTGGAC CCGGCCTGGAC CCGGCCTGGAC CCGGCCCGCA CCGTGGGCGC GCGCGACCGT TGCAGCCGCC 12801 CGTGGAGGAA TATGACGAGG ACGATGAGTA CGAGCCAGAG GACGGCGACT 12851 ACTAAGCGGT GATGTTTCTG ATCAGATGAT GCAAGACGCA ACGTCGCCGC 12901 CGGTGCGGGC GGCGCTGCAG AGCCAGCAG GACGGCCGCC 12901 CGGTGCGGGC GGCGCTGCAG AGCCAGCAG GGCCAGAG CCCGCCCGCCCC CCGCACCGT TGCAGCCCGC CGCCACCGC CCGCACCGT TGCAGCCCGC CCGCACCGC CCGCACCGT TGCAGCCGC CCGCACCGC CCGCACCGC CCGCACCGT CCGGCCTTAA CTCCACGGAC GCCACCCCG CCGCACCGT TGCGGCCCGACGT TGCGTCGGCC CCGCACCTC TGCGCCCTAA GCCAGCCGC CCGCACCTC TCGGTCGGCA GGCCGGAATT GAGGTGCCTG CCGCCACCGC CCGCACCGC CCGCACCGC CCGCACCAC CCGCCACCGC CCGCACCGC CCGCACCGC CCGCACCGC CCGCACCCC CCGCCACCCC CCGCCCCC CCGCCCCC CCGCCCCCC CCGCCCCC CCTTTGGGCC GCCCGAGAACCCCA CCGCCAGACAA CCGCCCACCAC CCGCCCCCC CCTTTAGACCC CCACACACAC CCGCCCACACCCC CCGCCCCC CCTTTTGGGGT GGCCGAGAA GGTGCTGCGC CCTTAAGACC CCACACCAC CCGCCCCCC CCTTTTGGGGT GCCCGAGACCCA CCGCCGCCCCCCCCCC	12001	ACCCCCAAAT	AGCGTTGCTC	GCGTAGGTGT	TCCGGCACTC	GCACTCGGCC
GCCGCGCTCG AGTCGCTGCC GCTCGACTAC GTGTCGGACG TTTCCCGGGA 12651 GGCTGGCACG GGCAGCGGCC ATAGAGAGGC CGAGTCCTAC TTTGACGCGG CCGACCGTGC CCGTCGCCGC TATCTCTCCG GCTCAGGATG AAACTGCGCC 12701 GCGCTGACCT GCGCTGGGCC CCAAGCCGAC GCGCCCTGGA GGCAGCTGGG CGCGCACTGGA CGCGACCCGG GGTTCGGCT CGCGGGACCT CCGTCGACCC 12751 GCCGGACTGG GGCTGGCGGT GGCACCCGCG CGCGCGGACCT CCGTCGACCC 12801 CGTGGAGGAA TATGACGAGG ACGATGAGTA CGAGCCAGAG GACGCCGGC CCGCCCCCCC CGCGCACCGCT TGCAGCCGCC 12801 CGTGGAGGAA TATGACGAGG ACGATGAGTA CGAGCCAGAG GACGCCGGCC 12851 ACTAAGCGGT GATGTTTCTG ATCAGATGAT GCAGACCAGA ACGGACCCGC TGCACCCCCC CTACAAAGAC TAGTCTACTA CGTTCTGCGT TGCCTGGCCC 12901 CGGTGCGGGC GGCGCTGCAG AGCCAGCCGT CCGCCGCCTAA CTCCACGGAC GCCACGCCCG CCGCGACGCT TCGGTCGCCT TCGGTCGCCT TGCCTGGCCCT CTGACCGCCC CCGCGACGCT TCGGTCGCA GGCCGGAATT GAGGTCCTG CTGACCGCCG CCGCGACGCT TCGGTCGCAC GGCCGGAATT GAGGTCCTG TCCAGCACGC CCGCGACGCC CCGCGACGCC CCGCGACGCC CCGCGACGCC CCGCGACGCC CCGCGACGCC CCGCGACGCC CCGCGACGCC CCGCGACGCC CCGCAGCACA CCGCCGCAATCC CGCCCGAACCCC CCGCCAACCCC CCGCACGACGC CCGCAGCACA CCGCCCGAACCCC CCGCACGACGC CCGCAGCACA CCGCCCGAACCCC CCGCACGACG CCGCCCGACGCC CCGCAGCACA CCGCCCGACGCC CCGCACGACGC CCGCCACGACGC CCGCCACGACGC CCGCCCGACGCC CCGCCGCCC CCGCCCGC						
12651 GGCTGGCACG GGCAGCGGCG ATAGAAGGC CGAGTCCTAC TTTGACGCGC CCGACCGTC CCGTCGCCCC TATCTCTCCG GCTCAGGATG AAACTGCGCC CCGACCGGC CCGACCGAC GCGCACCGGC GCGCGACCGC CCGCGACCGC CCGCGACCGC CCGCGACCGC CCGCGACCGC CCGCGACCGC CCGCGGACCT CCGTCGACCC CCGCGCCTGGAC CCGCGCCCGCG CGCGCGCCC CCGTCGACCC CCGCGCCCGC CCGCGCACCGC CCGCGCGCCC CCGCCGCCC CCGCCGCCC CCGCCG	12601	CGGCGCGAGC	TCAGCGACCG	CGAGCTGATG	CACAGCCTGC	MAAGGGCCC 1
CCGACCGTCC CCGTCGCCGC TATCTCTCCG GCTCAGGATG AAACTGCGCC 12701 GCGCTGACCT GCGCTGGGCC CCAAGCCGAC GCGCCCTGGA GGCAGCTGGG CGCGACTGGA CGCGACCCGG GGTTCGGCTG CGCGGACCTC CCGTCGACCC 12751 GCCGGACCTG GGCTGGCGGT GGCACCCGCG CGCGCGGACCT CCGTCGACCC 12801 CGTGGAGGAA TATGACGAGG ACGATGAGTA CGAGCCAGAG GACGCGAGT GCACCTCCTT ATACTGCTCC TGCTACTCAT GCTCGGTCTC CTGCCGCTCA 12851 ACTAAGCGGT GATGTTTCTG ATCAGATGAT GCAAGACGCA ACGGACCAGAG GACGCAGAG TAGATTCGCCA CTACAAAGAC TAGTCTACTA CGTTCTGCGT TGCCTGGCCC 12901 CGGTGCGGGC GGCGCTGCAG AGCCAGCCGT CCGGCCTTAA CTCCACGGAC GCCACGCCG CCGCGACGTC TCGGTCGGCA GGCCAGCCCG CCGCGACGTC TCGGTCGGCA GGCCGGAATT GAGGTGCCTG CTGACCGCGG TCCAGTACCT GGCGTAGTAC AGCGACTGAC GCCGCGTTAGC CCGCGCAATCC CTGACCGCGG TCCAGTACCT GGCGTAGTAC AGCGACTGAC GCCGCTTAGGAC GCCGTTAGGAC GCCGTAGAG GCCGCTTAGGAC CCGCAGAGAC CCGCAGAGAC CGCAATCCC ACGCCGTTA CTCCACGGAC GCCGTTAGGAC CGCAACACCA CCGAACAGAA GGTGCTGGC TTCGCCCACCA GGGCCGCGC CGTTTGGGGT GCCTGACAGAA GGTGCTGGCG CGTTTGGCCGC CGTTTGGGGT GCGTGCTCTT CCACGACCGC 13101 ATCGTAAACG CGCGCGCG GCAAACCCCA CGCACGAGAA GGTGCTGGCG TAGCCACCGC TTTGGCCCGC TTTTGTCCCGG TGCTCCGGCC ACGAGGCCGC TCCTCCGGCC TTTTGTCCCGG TGCTCCTGCCGCC TTTTGTCCCGG TGCTCCTTAC AACAGCGCCA AACACCCA CCGAGCAATG TTCTCCGGCCC TGCTCCGGCC TTTTGTCCCGG TGCTCCGGCC TGCTCCGGCC TTTTGTCCCGG TGCCCGGCC TGCTCCGGCC TTTTGTCCCGG TGCTCCGGCC TGCTCCGGCC TTTTGTCCCGG TGCCCGGCC TGCTCCGGCC TTTTGTCCCGG TGCCCCGGC TGCTCCGGCC TTTTGCCCGGC TGCTCCGGCC TGCTCCGGCC TTTTGTCCCGG CCGAGCCAATG TTTGTCCCGGC TGCTCCGGCC TGCTCCGGCC TGCTCCGGCC TGCTCCGGCC TGCCCGGCC TGCCCGGCCGCC CGGCCGCGC GGGCCGCGC GGGCCGCGC CCGAGCAATG TTGCCCCGCCCC						
12701 GCGCTGACCT GCGCTGGGCC CCAAGCCGAC GCGCCCTGGA GGCAGCTGGC CGCGGACTGGA CGCGACCCGG GGTTCGGCTG CGCGGGACCT CCGTCGACCC CGCGCACCTGGACCC CGCGCACCTGGACCC CGGCCTGGACC CCGCCGCACCCGC CGCGCACCGC CGGCCTGGAC CCGACCGCCA CCGTGGGCGC CGCGCACCGT TGCAGCCGCC CGGCCTGGAC CCGACCGCCA CCGTGGGCGC GCGCGACCGT TGCAGCCGCC CGGCCTGGAC CCGACCGCCA CCGTGGGCGC GCGCGACCGT TGCAGCCGCC CGGCCTCATA ATACTGCTCC TGCTACTCAT GCTCGGTCTC CTGCCGCTCA ACGTAGCGAC GACGCAGAG GACGGCGAGT TGATTCGCCA CTACAAAGAC TAGTCTACTA CGTTCTGCGT TGCCTGGGCC GCCACGCCG CCGCGACGTC TCGGTCGCA GGCCGGACGT TGCCTGGGCC CCGCGACGTC TCGGTCGGCA GGCCGGAATT GAGGTGCCTG CCGCCACGCCG CCGCGACGTC TCGGTCGGCA GGCCGGAATT GAGGTGCCTG TCCAGTACCT GGCGCAATCC CTGACCGCGG TCCAGTACCT GGCGTAGTAC AGCGACTGAC GCGCGTTAGG CCGCGCAATCC CGGCACGCAG GCCGCACGCCG CGCAGCAGA CCGGCTTAGG CGCGCTTAGGCC TTCGGCGCACGC CGCAGCCGA CCGGCTCTCC GCAATTCTGG ACTGCGCACAC GCCGCTCGTCG GCGTCGTCG GCGTTAAGACC TTCGCCACCAC GCCGCCGCCGC CGTTTGGGGT GCCTGGCGC CGTTAAGACCC ACCGCCGCCGC CGTTTAGGCCCCA CGCACGAGAA GGTGCTGGCG TTCGCCACCAC GGCCCGCCGC CGTTTGGGGT GCCTGGCCG TTCGCCACCAC GCGCCCGCC CGTTTGGGGT GCGTGCTCTC CCACGACCGC TTCGCCACCAC GCGCCCGCC TTTCGCCCCG ACGAGCCGC TGCTCCGCCC ACGAGCCCGC TTCCCGCCC TTCGCCCCG ACGAGCCGC TGCTCCGCCC ACGAGCCCGC TGCTCCCGCCC TTCGCCCCGCC TTCCCGCCC TTCCCGCCCC AACCCCCA CCGAGCAATG TTCTCCCGCCT TTCTCCCGCCCC TTCCCGCCCC TTCCCCGCCC TTCCCGCCCC TTCCCCCCCC	12651	GGCTGGCACG	GGCAGCGGCG	ATAGAGAGGC	CGAGTCCTAC	TTTGACGCGG
CGCGACTGGA CGCGACCCGG GGTTCGGCTG CGCGGGACCT CCGTCGACCC 12751 GCCGGACCTG GGCTGGCGGT GGCACCCGCG CGCGCTGGCA ACGTCGGCGG CGGCCTGGAC CCGACCGCCA CCGTGGGCGC GCGCGACCGT TGCAGCCGCC CCGTGGGCGC GCGCGACCGT TGCAGCCGCC CCGTGGGCGC GCGCGACCGT TGCAGCCGCC CCGGCCTGGAC CCGACCGCC CCGACCGCC CCGACCGCC TGCAGCCAGAG GACGGCGAGT GCACCTCCTT ATACTGCTCC TGCTACTCAT GCTCGGTCTC CTGCCGCTCA ATACTGCTCC TGCTACTCAT GCTCGGTCTC CTGCCGCTCA TGATTCGCCA CTACAAAGAC TAGTCTACTA CGTTCTGCGT TGCCTGGGCC GCCACGCCGG CCGCGACCGC CCGCGACCGT CCGGCCTTAA CTCCACGGAC GCCACGCCCG CCGCGACCGC CCGCGACCTC TCGGTCGGCA GGCCGGAATT GAGGTGCCTG CTGACCGCG CCGCGACCGC CCGCATCATG TCGCTGACTG CGCCGAATCC CTGACCGCGG TCCAGTACCT GGCGACCTGAC GCGCCGTTAGG CCGCCGTTAGG CCGCCGTTAGG CCGCGCAATCC CGCACCGCG TCCAGTACCT GCCGACCGAC CCGCACTCC GCAATTCTGG ACTGCGCAAG GCCGTCGTCG GCGTCCGGTT GGCCGAAGAG CCGTTAAGACC TTCGCCGCAAG GCCGTTCGGCG CGTTAAGACC CCGACGAGAA GGTGCTGGCG CGTTCGCCGCG CCGTTGGGGT GCCGAGAGA GGTGCTGGCG TTCGCCACCA GGGCCGCG CCGTTTGGGGT GCCGAGAACCCCA CGCACGAGAA GGTGCTGGCG TTCGCCACCA GGGCCGCGC CGTTTGGGGT GCGTGCTCTT CCACGACCGC TTCGCCACCAC CGCACGAGAA GGTGCTGGCG TTCGCCGCCG TTCGCCGCCG TTCGCCGCC TTCGCCGCC TTCGCCGCCC TTCGCCGCCC TTCGCCGCCC TTCGCCCGCC		CCGACCGTGC	CCGTCGCCGC	TATCTCTCCG	GCTCAGGATG	AAACTGCGCC
CGCGACTGGA CGCGACCCGG GGTTCGGCTG CGCGGGACCT CCGTCGACCC 12751 GCCGGACCTG GGCTGGCGGT GGCACCCGCG CGCGCTGGCA ACGTCGGCGG CGGCCTGGAC CCGACCGCCA CCGTGGGCGC GCGCGACCGT TGCAGCCGCC CCGTGGGCGC GCGCGACCGT TGCAGCCGCC CCGTGGGCGC GCGCGACCGT TGCAGCCGCC CCGGCCTGGAC CCGACCGCC CCGACCGCC CCGACCGCC TGCAGCCAGAG GACGGCGAGT GCACCTCCTT ATACTGCTCC TGCTACTCAT GCTCGGTCTC CTGCCGCTCA ATACTGCTCC TGCTACTCAT GCTCGGTCTC CTGCCGCTCA TGATTCGCCA CTACAAAGAC TAGTCTACTA CGTTCTGCGT TGCCTGGGCC GCCACGCCGG CCGCGACCGC CCGCGACCGT CCGGCCTTAA CTCCACGGAC GCCACGCCCG CCGCGACCGC CCGCGACCTC TCGGTCGGCA GGCCGGAATT GAGGTGCCTG CTGACCGCG CCGCGACCGC CCGCATCATG TCGCTGACTG CGCCGAATCC CTGACCGCGG TCCAGTACCT GGCGACCTGAC GCGCCGTTAGG CCGCCGTTAGG CCGCCGTTAGG CCGCGCAATCC CGCACCGCG TCCAGTACCT GCCGACCGAC CCGCACTCC GCAATTCTGG ACTGCGCAAG GCCGTCGTCG GCGTCCGGTT GGCCGAAGAG CCGTTAAGACC TTCGCCGCAAG GCCGTTCGGCG CGTTAAGACC CCGACGAGAA GGTGCTGGCG CGTTCGCCGCG CCGTTGGGGT GCCGAGAGA GGTGCTGGCG TTCGCCACCA GGGCCGCG CCGTTTGGGGT GCCGAGAACCCCA CGCACGAGAA GGTGCTGGCG TTCGCCACCA GGGCCGCGC CGTTTGGGGT GCGTGCTCTT CCACGACCGC TTCGCCACCAC CGCACGAGAA GGTGCTGGCG TTCGCCGCCG TTCGCCGCCG TTCGCCGCC TTCGCCGCC TTCGCCGCCC TTCGCCGCCC TTCGCCGCCC TTCGCCCGCC	12701	GCGCTGACCT	GCGCTGGGCC	CCAAGCCGAC	GCGCCCTGGA	GGCAGCTGGG
12751 GCCGGACCTG GGCTGGCGGT GGCACCCGCG CGCGCTGGCA ACGTCGGCGC CGGCCTGGAC CCGACCGCCA CCGTGGGCGC GCGCGACCGT TGCAGCCGCC CCGACCGT TGCAGCCGCC CCGACCGT TGCAGCCGCC CCGACCGT TGCAGCCGCC GCGCGACCGT TGCAGCCGCC GCGCCGACCGT TGCAGCCGCC TATACCTGCTCC TGCTACTCAT GCTCGGTCTC CTGCCGCTCA ATACTGCTCC TGCTACTCAT GCTCGGTCTC CTGCCGCTCA ACGTACCAAAGAC TAGTCTACTA CGAAGACGCA ACGGACCCGG TGATTCGCCA CTACAAAGAC TAGTCTACTA CGTTCTGCGT TGCCTGGGCC GCCACGCCCG CCGCGACCGT TCGGTCGGCA GGCCGGAATT GAGGTGCCTG CCGCCACGCCC CCGCGACCGT TCGGTCGGCA GGCCGGAATT GAGGTGCCTG CTGACCGGGC TCCAGTACCT GGCGACTGAC GCCGCAATCC CTGACCGCGG TCCAGTACCT GGCGTAGTAC AGCGACTGAC GCGCGTTAGG CCGCAATCC GCGCAATCC GCGCAGACG GCCGCAATCC GCGCAGACG GCCGCAATCC GCGCAGAACCCA ACGCCGAATCC GCGCTTAAACCC GCCGCCACGC CGTTAAGACC TTCGCCGCAA CCGGCTCTCC GCAATTCTGG GCCGCAACCCA GGGCCGAAGA GGTGCTGGCG TTCGCCGCACAG GCCGTTCGGGT GCCGAGAAGA GGTGCTGGCG TTCGCCACCA GGGCCGCGC CGTTTGGGGT GCGTGCTCT CCACGACCGC TTCGCCACCA GGGCCGCGC CGTTTGGGGT GCGTGCTCTT CCACGACCGC TTCGCCACCA GGGCCGGCC TTCGCCGCCG TTCGCGCCG TTCGCCGCC TTCGGCCCG TAGGCCCGG TTCCCGGCCG TTCCCGGCCC TTCCGGCCCG TAGGCCGGC TTCCCGGCCG TTCCCGGCCCG TTCCCGGCCCG TTCCCGGCCCG TTCCCGGCCCG TTCCCGGCCCG TTCCCGGCCCT TTCGCCGCCC TAGGCCGGCC TTCCCGGCCCT TTCGCCGCCC TTCCCGCCCC TTCCCGCCCCT TTCGCCGCCC TTCCCGCCCCT TTCCCGCCCCT TTCCCGCCCCT TTCCCGCCCT TTCCCGCCCCT TTCCCGCCCT TTCCCCGCCCT TTCCCCGCCCT TTCCCCGCCCT TTCCCCGCCCT TTCCCGCCCT TTCCCCGCCCT TTCCCCGCCCT TTCCCGCCCT TTCCCCCCCT TTCCCCCCCT TTCCCCCCCT TTCCCCCC						
CGGCCTGGAC CCGACCGCA CCGTGGGCGC GCGCGACCGT TGCAGCCGCC 12801 CGTGGAGGAA TATGACGAGG ACGATGAGTA CGAGCCAGAG GACGGCGAGT GCACCTCCTT ATACTGCTCC TGCTACTCAT GCTCGGTCTC CTGCCGCTCA 12851 ACTAAGCGGT GATGTTTCTG ATCAGATGAT GCAAGACGCA ACGGACCCGG TGATTCGCCA CTACAAAGAC TAGTCTACTA CGTTCTGCGT TGCCTGGGCC 12901 CGGTGCGGGC GGCGCTGCAG AGCCAGCCGT CCGGCCTTAA CTCCACGGAC GCCACGCCCG CCGCGACGTC TCGGTCGGCA GGCCGGAATT GAGGTGCCTG 12951 GACTGGCGCC AGGTCATGGA CCGCATCATG TCGCTGACTG CGCGCAATCC CTGACCGCG TCCAGTACCT GGCGTAGTAC AGCGACTGAC GCGCTTAGG 13001 TGACGCGTTC CGGCAGCAGC CGCAGGCCAA CCGGCTCTCC GCAATTCTGG ACTGCGCAAG GCCGTCGTCG GCGTCCGGTT GGCCGAGAGG CGTTAAGACC 13051 AAGCGGTGGT CCCGGCCGC GCAAACCCCA CGCACGAGAA GGTGCTGGCG TTCGCCACCAC GGCCCGCGC CGTTTGGGGT GCGTGCTCT CCACGACCGC 13101 ATCGTAAACG CGCTGGCCGA AAACAGGGCC ATCCGGCCCG ACGAGCCGC 13151 CCTGGTCTAC GACGCGCTGC TTCAGCGCGT TAGGCCGGCC TGCTCCGGCC 13151 CCTGGTCTAC GACGCGCTGC TTCAGCGCGT GGCTCGTTAC AACAGCGGCA 13201 ACGTGCAGAC CAACCTGGAC CGGCTGGTGG GGGATGTGCG CGGAGCCGTG 13201 ACGTGCAGAC CAACCTGGAC CGGCTGGTGG GGGATGTGCG CGAGGCCGTG						
CGTGGAGGAA TATGACGAGG ACGATGAGTA CGAGCCAGAG GACGGCGAGT GCACCTCCTT ATACTGCTCC TGCTACTCAT GCTCGGTCTC CTGCCGCTCA 12851 ACTAAGCGGT GATGTTTCTG ATCAGATGAT GCAAGACGCA ACGGACCCGG TGATTCGCCA CTACAAAGAC TAGTCTACTA CGTTCTGCGT TGCCTGGGCC 12901 CGGTGCGGGC GGCGCTGCAG AGCCAGCCGT CCGGCCTTAA CTCCACGGAC GCCACGCCCG CCGCGACGTC TCGGTCGGCA GGCCGGAATT GAGGTGCCTG CTGACCGGC AGGTCATGA CTCCACGGAC GCCACGCCCG TCCAGTACCT GGCGTAGTAC AGCGACTGAC GCGCGTTAGG 12951 GACTGGCGCC AGGTCATGGA CCGCATCATG TCGCTGACTG CGCGCAATCC CTGACCGGG TCCAGTACCT GGCGTAGTAC AGCGACTGAC GCGCGTTAGG 13001 TGACGCGTTC CGGCAGCAGC CGCAGGCCAA CCGGCTCTCC GCAATTCTGG ACTGCGCAAG GCCGTCGTCG GCGTCCGGTT GGCCGAAGAG CGTTAAGACC 13051 AAGCGGTGGT CCCGGCGCG GCAAACCCCA CGCACGAGAA GGTGCTGGCG TTCGCCACCA GGGCCGCGC CGTTTGGGGT GCGTGCTCTT CCACGACCGC 13101 ATCGTAAACG CGCTGGCCGA AAACAGGGCC ATCCGGCCCG ACGAGGCCGG TAGCCATTTCC GCGACCGCC TTCGCCGGCC TTCGCCGGCC TTCGCCGGCC TTCGCCGGCC TTCGCCGGCC TTCTCCGGCCCG TTCCCGGCCCGCC TTCGCCGGCC TTCGCCGGCC TTCTCCGGCCCGCCGCCGCGCGCG	12751	GCCGGACCTG	GGCTGGCGGT	GGCACCCGCG	CGCGCTGGCA	ACGTCGGCGG
GCACCTCCTT ATACTGCTCC TGCTACTCAT GCTCGGTCTC CTGCCGCTCA 12851 ACTAAGCGGT GATGTTTCTG ATCAGATGAT GCAAGACGCA ACGGACCCGG TGATTCGCCA CTACAAAGAC TAGTCTACTA CGTTCTGCGT TGCCTGGGCC 12901 CGGTGCGGGC GGCGCTGCAG AGCCAGCCGT CCGGCCTTAA CTCCACGGAC GCCACGCCCG CCGCGACGTC TCGGTCGGCA GGCCGGAATT GAGGTGCCTG 12951 GACTGGCGCC AGGTCATGGA CCGCATCATG TCGCTGACTG CGCGCAATCC CTGACCGCGG TCCAGTACCT GGCGTAGTAC AGCGACTGAC GCGCGTTAGG 13001 TGACGCGTTC CGGCAGCAGC CGCAGGCCAA CCGGCTCTCC GCAATTCTGG ACTGCGCAAG GCCGTCGTCG GCGTCCGGTT GGCCGAGAGA GGTGCTGGCG TTCGCCACCA GGGCCGCGC CGTTTGGGGT GCCTGACAGA GGTGCTGGCG 13051 AAGCGGTGGT CCCGGCGCGC CGTTTGGGGT GCGTGCTCTT CCACGACCGC 13101 ATCGTAAACG CGCTGGCCGA AAACAGGGCC ATCCGGCCCG ACGAGCCGC TAGCATTTTGC GCGACCGGCT TTTGTCCCGG TAGGCCGGC TGCTCCGGCC 13151 CCTGGTCTAC GACGCGCTGC TTCAGCGCGT GGCTCGTTAC AACAGCGCCA GGACCAGATG CTGCCGGAC CCGAGCAATG TTGTCGCCGT 13201 ACGTGCAGAC CAACCTGGAC CGGCTGGTGG GGGATGTGCG CGAGGCCGGT 13201 ACGTGCAGAC CAACCTGGAC CGGCTGGTGG GGGATGTGCG CGAGGCCGGT		CGGCCTGGAC	CCGACCGCCA	CCGTGGGCGC	GCGCGACCGT	TGCAGCCGCC
GCACCTCCTT ATACTGCTCC TGCTACTCAT GCTCGGTCTC CTGCCGCTCA 12851 ACTAAGCGGT GATGTTTCTG ATCAGATGAT GCAAGACGCA ACGGACCCGG TGATTCGCCA CTACAAAGAC TAGTCTACTA CGTTCTGCGT TGCCTGGGCC 12901 CGGTGCGGGC GGCGCTGCAG AGCCAGCCGT CCGGCCTTAA CTCCACGGAC GCCACGCCCG CCGCGACGTC TCGGTCGGCA GGCCGGAATT GAGGTGCCTG 12951 GACTGGCGCC AGGTCATGGA CCGCATCATG TCGCTGACTG CGCGCAATCC CTGACCGCGG TCCAGTACCT GGCGTAGTAC AGCGACTGAC GCGCGTTAGG 13001 TGACGCGTTC CGGCAGCAGC CGCAGGCCAA CCGGCTCTCC GCAATTCTGG ACTGCGCAAG GCCGTCGTCG GCGTCCGGTT GGCCGAGAGA GGTGCTGGCG TTCGCCACCA GGGCCGCGC CGTTTGGGGT GCCTGACAGA GGTGCTGGCG 13051 AAGCGGTGGT CCCGGCGCGC CGTTTGGGGT GCGTGCTCTT CCACGACCGC 13101 ATCGTAAACG CGCTGGCCGA AAACAGGGCC ATCCGGCCCG ACGAGCCGC TAGCATTTTGC GCGACCGGCT TTTGTCCCGG TAGGCCGGC TGCTCCGGCC 13151 CCTGGTCTAC GACGCGCTGC TTCAGCGCGT GGCTCGTTAC AACAGCGCCA GGACCAGATG CTGCCGGAC CCGAGCAATG TTGTCGCCGT 13201 ACGTGCAGAC CAACCTGGAC CGGCTGGTGG GGGATGTGCG CGAGGCCGGT 13201 ACGTGCAGAC CAACCTGGAC CGGCTGGTGG GGGATGTGCG CGAGGCCGGT	12801	CGTGGAGGAA	TATGACGAGG	ACGATGAGTA	CGAGCCAGAG	GACGGCGAGT
TGATTCGCCA CTACAAAGAC TAGTCTACTA CGTTCTGCGT TGCCTGGGCC 12901 CGGTGCGGGC GGCGCTGCAG AGCCAGCCGT CCGGCCTTAA CTCCACGGAC GCCACGCCCG CCGCGACGTC TCGGTCGGCA GGCCGGAATT GAGGTGCCTG 12951 GACTGGCGCC AGGTCATGGA CCGCATCATG TCGCTGACTG CGCGCAATCC CTGACCGCGG TCCAGTACCT GGCGTAGTAC AGCGACTGAC GCGCGTTAGG 13001 TGACGCGTTC CGGCAGCAGC CGCAGGCCAA CCGGCTCTCC GCAATTCTGG ACTGCGCAAG GCCGTCGTCG GCGTCCGGTT GGCCGAGAGG CGTTAAGACC 13051 AAGCGGTGGT CCCGGCGCGC GCAAACCCCA CGCACGAGAA GGTGCTGGCG TCCGCCACCA GGGCCGCGCG CGTTTGGGGT GCGTGCTCTT CCACGACCGC 13101 ATCGTAAACG CGCTGGCCGA AAACAGGGCC ATCCGGCCCG ACGAGGCCGGC TAGCATTTGC GCGACCGGCT TTTTGTCCCGG TAGGCCGGGC TGCTCCGGCCC TGCTCCGGCCC AACACCCCA CGCACGAATG TTTTCCCCGGCCAAACCCCA CCGAGCAATG TTTTCCCCGGT AACAGCCGCAAACCCCA CCGAGCAATG TTTTCCCCGGCCAAACCCCAAACCCCAAACCCCAACCGCCAACCGCCAAACCCCAACCGCCAACCGCCAAACCCCAACCGCCAACCGCCAACCGCCAACCGCCAACCGCCAACCGCCCAACCGCCCAACCGCCCAACCGCCCAAACCCCAACCGCCCAACCCGCCCAACCCGCCCAACCCGCCCAACCCGCCCAACCCGCCCAACCCGCCCAACCCGCCCAACCCGCCCAACCCGCCCAACCCGCCCAACCCGCCCAACCCGCCCAACCCGCCAACCCGCCCAACCCGCCCAACCCGCCAACCCCCAACCCGCCAACCCGCCAACCCCCAACCCGCCAACCCGCCAACCCCCAACCCCCAACCCCCAACCCCCC	12001	GCACCTCCTT	ATACTGCTCC	TGCTACTCAT	GCTCGGTCTC	CTGCCGCTCA
TGATTCGCCA CTACAAAGAC TAGTCTACTA CGTTCTGCGT TGCCTGGGCC 12901 CGGTGCGGGC GGCGCTGCAG AGCCAGCCGT CCGGCCTTAA CTCCACGGAC GCCACGCCCG CCGCGACGTC TCGGTCGGCA GGCCGGAATT GAGGTGCCTG 12951 GACTGGCGCC AGGTCATGGA CCGCATCATG TCGCTGACTG CGCGCAATCC CTGACCGCGG TCCAGTACCT GGCGTAGTAC AGCGACTGAC GCGCGTTAGG 13001 TGACGCGTTC CGGCAGCAGC CGCAGGCCAA CCGGCTCTCC GCAATTCTGG ACTGCGCAAG GCCGTCGTCG GCGTCCGGTT GGCCGAGAGG CGTTAAGACC 13051 AAGCGGTGGT CCCGGCGCGC GCAAACCCCA CGCACGAGAA GGTGCTGGCG TCCGCCACCA GGGCCGCGCG CGTTTGGGGT GCGTGCTCTT CCACGACCGC 13101 ATCGTAAACG CGCTGGCCGA AAACAGGGCC ATCCGGCCCG ACGAGGCCGGC TAGCATTTGC GCGACCGGCT TTTTGTCCCGG TAGGCCGGGC TGCTCCGGCCC TGCTCCGGCCC AACACCCCA CGCACGAATG TTTTCCCCGGCCAAACCCCA CCGAGCAATG TTTTCCCCGGT AACAGCCGCAAACCCCA CCGAGCAATG TTTTCCCCGGCCAAACCCCAAACCCCAAACCCCAACCGCCAACCGCCAAACCCCAACCGCCAACCGCCAAACCCCAACCGCCAACCGCCAACCGCCAACCGCCAACCGCCAACCGCCCAACCGCCCAACCGCCCAACCGCCCAAACCCCAACCGCCCAACCCGCCCAACCCGCCCAACCCGCCCAACCCGCCCAACCCGCCCAACCCGCCCAACCCGCCCAACCCGCCCAACCCGCCCAACCCGCCCAACCCGCCCAACCCGCCCAACCCGCCAACCCGCCCAACCCGCCCAACCCGCCAACCCCCAACCCGCCAACCCGCCAACCCCCAACCCGCCAACCCGCCAACCCCCAACCCCCAACCCCCAACCCCCC	12851	ACTAAGCGGT	GATGTTTCTG	ATCAGATGAT	GCAAGACGCA	ACGGACCCGG
12901 CGGTGCGGC GGCGCTGCAG AGCCAGCCGT CCGGCCTTAA CTCCACGGAC GCCACGCCCG CCGCGACGTC TCGGTCGGCA GGCCGGAATT GAGGTGCCTG 12951 GACTGGCGCC AGGTCATGGA CCGCATCATG TCGCTGACTG CGCGCAATCC CTGACCGCGG TCCAGTACCT GGCGTAGTAC AGCGACTGAC GCGCGTTAGG 13001 TGACGCGTTC CGGCAGCAGC CGCAGGCCAA CCGGCTCTCC GCAATTCTGG ACTGCCGAAG GCCGTCGTCG GCGTCCGGTT GGCCGAGAGC CGTTAAGACC 13051 AAGCGGTGGT CCCGGCGCGC GCAAACCCCA CGCACGAGAA GGTGCTGGCG TCCGCCACCA GGGCCGCGCG CGTTTGGGGT GCGTGCTCTT CCACGACCGC 13101 ATCGTAAACG CGCTGGCCGA AAACAGGGCC ATCCGGCCCG ACGAGGCCGGC TAGCATTTTGC GCGACCGGCT TTTGTCCCGG TAGGCCGGGC TGCTCCGGCC 13151 CCTGGTCTAC GACGCGCTGC TTCAGCGCGT GGCTCGTTAC AACAGCGGCA GGACCAGATG CTGCGCGGAAGCCGGC AAGTCGCGCA CCGAGCAATG TTGTCGCCGT 13201 ACGTGCAGAC CAACCTGGAC CGGCTGGTGG GGGATGTGCG CGAGGCCGTG		TGATTCGCCA	CTACAAAGAC	TAGTCTACTA	CGTTCTGCGT	TGCCTGGGCC
GCCACGCCG CCGCGACGTC TCGGTCGGCA GGCCGGAATT GAGGTGCCTG 12951 GACTGGCGCC AGGTCATGGA CCGCATCATG TCGCTGACTG CGCGCAATCC CTGACCGCGG TCCAGTACCT GGCGTAGTAC AGCGACTGAC GCGCGTTAGG 13001 TGACGCGTTC CGGCAGCAGC CGCAGGCCAA CCGGCTCTCC GCAATTCTGG ACTGCGCAAG GCCGTCGTCG GCGTCCGGTT GGCCGAGAGG CGTTAAGACC 13051 AAGCGGTGGT CCCGGCGCGC GCAAACCCCA CGCACGAGAA GGTGCTGGCG TTCGCCACCA GGGCCGCGC CGTTTGGGGT GCGTGCTCTT CCACGACCGC 13101 ATCGTAAACG CGCTGGCCGA AAACAGGGCC ATCCGGCCCG ACGAGGCCGG TAGCATTTGC GCGACCGGCT TTTGTCCCGG TAGGCCGGGC TGCTCCGGCC 13151 CCTGGTCTAC GACGCGCTGC TTCAGCGCGT GGCTCGTTAC AACAGCGGCA GGACCAGATG CTGCGCGACC AAGTCGCGCA CCGAGCAATG TTGTCGCCGT 13201 ACGTGCAGAC CAACCTGGAC CGGCTGGTGG GGGATGTGCG CGAGGCCGTG						
GACTGGCGCC AGGTCATGGA CCGCATCATG TCGCTGACTG CGCGCAATCC CTGACCGCGG TCCAGTACCT GGCGTAGTAC AGCGACTGAC GCGCGTTAGG 13001 TGACGCGTTC CGGCAGCAGC CGCAGGCCAA CCGGCTCTCC GCAATTCTGG ACTGCGCAAG GCCGTCGTCG GCGTCCGGTT GGCCGAGAGAG CGTTAAGACC 13051 AAGCGGTGGT CCCGGCGCGC GCAAACCCCA CGCACGAGAA GGTGCTGGCG TCCGCCACCA GGGCCGCGC CGTTTGGGGT GCGTGCTCTT CCACGACCGC 13101 ATCGTAAACG CGCTGGCCGA AAACAGGGCC ATCCGGCCCG ACGAGGCCGG TAGCATTTGC GCGACCGGCT TTTGTCCCGG TAGGCCGGGC TGCTCCGGCC 13151 CCTGGTCTAC GACGCGCTGC TTCAGCGCGT GGCTCGTTAC AACAGCGGCA GGACCAGATG CTGCGCCGGT AAGTCGCGCA CCGAGCAATG TTGTCGCCGT 13201 ACGTGCAGAC CAACCTGGAC CGGCTGGTGG GGGATGTGCG CGAGGCCGTG	12901	CGGTGCGGGC	GGCGCTGCAG	AGCCAGCCGT	CCGGCCTTAA	CTCCACGGAC
CTGACCGCGG TCCAGTACCT GGCGTAGTAC AGCGACTGAC GCGCGTTAGG 13001 TGACGCGTTC CGGCAGCAGC CGCAGGCCAA CCGGCTCTCC GCAATTCTGG ACTGCGCAAG GCCGTCGTCG GCGTCCGGTT GGCCGAGAGG CGTTAAGACC 13051 AAGCGGTGGT CCCGGCGCGC GCAAACCCCA CGCACGAGAA GGTGCTGGCG TTCGCCACCA GGGCCGCGC CGTTTGGGGT GCGTGCTCTT CCACGACCGC 13101 ATCGTAAACG CGCTGGCCGA AAACAGGGCC ATCCGGCCCG ACGAGGCCGG TAGCATTTGC GCGACCGGCT TTTGTCCCGG TAGGCCGGGC TGCTCCGGCC 13151 CCTGGTCTAC GACGCGCTGC TTCAGCGCGT GGCTCGTTAC AACAGCGGCA GGACCAGATG CTGCGCGACC AAGTCGCGCA CCGAGCAATG TTGTCGCCGT 13201 ACGTGCAGAC CAACCTGGAC CGGCTGGTGG GGGATGTGCG CGAGGCCGTG						
CTGACCGCGG TCCAGTACCT GGCGTAGTAC AGCGACTGAC GCGCGTTAGG 13001 TGACGCGTTC CGGCAGCAGC CGCAGGCCAA CCGGCTCTCC GCAATTCTGG ACTGCGCAAG GCCGTCGTCG GCGTCCGGTT GGCCGAGAGG CGTTAAGACC 13051 AAGCGGTGGT CCCGGCGCGC GCAAACCCCA CGCACGAGAA GGTGCTGGCG TTCGCCACCA GGGCCGCGC CGTTTGGGGT GCGTGCTCTT CCACGACCGC 13101 ATCGTAAACG CGCTGGCCGA AAACAGGGCC ATCCGGCCCG ACGAGGCCGG TAGCATTTGC GCGACCGGCT TTTGTCCCGG TAGGCCGGGC TGCTCCGGCC 13151 CCTGGTCTAC GACGCGCTGC TTCAGCGCGT GGCTCGTTAC AACAGCGGCA GGACCAGATG CTGCGCGACC AAGTCGCGCA CCGAGCAATG TTGTCGCCGT 13201 ACGTGCAGAC CAACCTGGAC CGGCTGGTGG GGGATGTGCG CGAGGCCGTG	12951	GACTGGCGCC	AGGTCATGGA	CCGCATCATG	TCGCTGACTG	CGCGCAATCC
ACTGCGCAAG GCCGTCGTCG GCGTCCGGTT GGCCGAGAGG CGTTAAGACC 13051 AAGCGGTGGT CCCGGCGCGC GCAAACCCCA CGCACGAGAA GGTGCTGGCG TTCGCCACCA GGGCCGCGC CGTTTGGGGT GCGTGCTCTT CCACGACCGC 13101 ATCGTAAACG CGCTGGCCGA AAACAGGGCC ATCCGGCCCG ACGAGGCCGG TAGCATTTGC GCGACCGGCT TTTGTCCCGG TAGGCCGGGC TGCTCCGGCC 13151 CCTGGTCTAC GACGCGCTGC TTCAGCGCGT GGCTCGTTAC AACAGCGGCA GGACCAGATG CTGCGCGACG AAGTCGCGCA CCGAGCAATG TTGTCGCCGT 13201 ACGTGCAGAC CAACCTGGAC CGGCTGGTGG GGGATGTGCG CGAGGCCGTG		CTGACCGCGG	TCCAGTACCT	GGCGTAGTAC	AGCGACTGAC	GCGCGTTAGG
ACTGCGCAAG GCCGTCGTCG GCGTCCGGTT GGCCGAGAGG CGTTAAGACC 13051 AAGCGGTGGT CCCGGCGCGC GCAAACCCCA CGCACGAGAA GGTGCTGGCG TTCGCCACCA GGGCCGCGC CGTTTGGGGT GCGTGCTCTT CCACGACCGC 13101 ATCGTAAACG CGCTGGCCGA AAACAGGGCC ATCCGGCCCG ACGAGGCCGG TAGCATTTGC GCGACCGGCT TTTGTCCCGG TAGGCCGGGC TGCTCCGGCC 13151 CCTGGTCTAC GACGCGCTGC TTCAGCGCGT GGCTCGTTAC AACAGCGGCA GGACCAGATG CTGCGCGACG AAGTCGCGCA CCGAGCAATG TTGTCGCCGT 13201 ACGTGCAGAC CAACCTGGAC CGGCTGGTGG GGGATGTGCG CGAGGCCGTG	13001	からな たらできずがた	CCCCACCACC	CGCAGGCCAA	CCGGCTCTCC	GCAATTCTGG
13051 AAGCGGTGGT CCCGGCGCGC GCAAACCCCA CGCACGAGAA GGTGCTGGCG TTCGCCACCA GGGCCGCGCG CGTTTGGGGT GCGTGCTCTT CCACGACCGC 13101 ATCGTAAACG CGCTGGCCGA AAACAGGGCC ATCCGGCCCG ACGAGGCCGG TAGCATTTGC GCGACCGGCT TTTGTCCCGG TAGGCCGGGC TGCTCCGGCC 13151 CCTGGTCTAC GACGCGCTGC TTCAGCGCGT GGCTCGTTAC AACAGCGGCA GGACCAGATG CTGCGCGACG AAGTCGCGCA CCGAGCAATG TTGTCGCCGT 13201 ACGTGCAGAC CAACCTGGAC CGGCTGGTGG GGGATGTGCG CGAGGCCGTG	13001	ACTGCGCAAG	CCCCTCCTCC	GCGTCCGGTT	GGCCGAGAGG	CGTTAAGACC
TTCGCCACCA GGGCCGCGC CGTTTGGGGT GCGTGCTCTT CCACGACCGC 13101 ATCGTAAACG CGCTGGCCGA AAACAGGGCC ATCCGGCCCG ACGAGGCCGG TAGCATTTGC GCGACCGGCT TTTGTCCCGG TAGGCCGGGC TGCTCCGGCC 13151 CCTGGTCTAC GACGCGCTGC TTCAGCGCGT GGCTCGTTAC AACAGCGGCA GGACCAGATG CTGCGCGACG AAGTCGCGCA CCGAGCAATG TTGTCGCCGT 13201 ACGTGCAGAC CAACCTGGAC CGGCTGGTGG GGGATGTGCG CGAGGCCGTG						
TTCGCCACCA GGGCCGCGC CGTTTGGGGT GCGTGCTCTT CCACGACCGC 13101 ATCGTAAACG CGCTGGCCGA AAACAGGGCC ATCCGGCCCG ACGAGGCCGG TAGCATTTGC GCGACCGGCT TTTGTCCCGG TAGGCCGGGC TGCTCCGGCC 13151 CCTGGTCTAC GACGCGCTGC TTCAGCGCGT GGCTCGTTAC AACAGCGGCA GGACCAGATG CTGCGCGACG AAGTCGCGCA CCGAGCAATG TTGTCGCCGT 13201 ACGTGCAGAC CAACCTGGAC CGGCTGGTGG GGGATGTGCG CGAGGCCGTG	13051	AAGCGGTGGT	CCCGGCGCGC	GCAAACCCCA	CGCACGAGAA	GGTGCTGGCG
TAGCATTTGC GCGACCGGCT TTTGTCCCGG TAGGCCGGGC TGCTCCGGCC 13151 CCTGGTCTAC GACGCGCTGC TTCAGCGCGT GGCTCGTTAC AACAGCGGCA GGACCAGATG CTGCGCGACG AAGTCGCGCA CCGAGCAATG TTGTCGCCGT 13201 ACGTGCAGAC CAACCTGGAC CGGCTGGTGG GGGATGTGCG CGAGGCCGTG		TTCGCCACCA	GGGCCGCGCG	CGTTTGGGGT	GCGTGCTCTT	CCACGACCGC
TAGCATTTGC GCGACCGGCT TTTGTCCCGG TAGGCCGGGC TGCTCCGGCC 13151 CCTGGTCTAC GACGCGCTGC TTCAGCGCGT GGCTCGTTAC AACAGCGGCA GGACCAGATG CTGCGCGACG AAGTCGCGCA CCGAGCAATG TTGTCGCCGT 13201 ACGTGCAGAC CAACCTGGAC CGGCTGGTGG GGGATGTGCG CGAGGCCGTG	12101	አመርርጥልልልርር	CCCTCCCCCA	AAACAGGGCC	ATCCGGCCCG	ACGAGGCCGG
13151 CCTGGTCTAC GACGCGCTGC TTCAGCGCGT GGCTCGTTAC AACAGCGGCA GGACCAGATG CTGCGCGACG AAGTCGCGCA CCGAGCAATG TTGTCGCCGT 13201 ACGTGCAGAC CAACCTGGAC CGGCTGGTGG GGGATGTGCG CGAGGCCGTG	13101	MICGIMANCG	CCCACCCCA		TAGGCCGGC	TGCTCCGGCC
GGACCAGATG CTGCGCGACG AAGTCGCGCA CCGAGCAATG TTGTCGCCGT 13201 ACGTGCAGAC CAACCTGGAC CGGCTGGTGG GGGATGTGCG CGAGGCCGTG		TAGCATTIGC	GCONCCOGC 1	111010000	1.10000000	
GGACCAGATG CTGCGCGACG AAGTCGCGCA CCGAGCAATG TTGTCGCCGT 13201 ACGTGCAGAC CAACCTGGAC CGGCTGGTGG GGGATGTGCG CGAGGCCGTG	13151	CCTGGTCTAC	GACGCGCTGC	TTCAGCGCGT	GGCTCGTTAC	AACAGCGGCA
13201 ACGTGCAGAC CAACCTGGAC CGGCTGGTGG GGGATGTGCG CGAGGCCGTG		GGACCAGATG	CTGCGCGACG	AAGTCGCGCA	CCGAGCAATG	TTGTCGCCGT
13201 ACGTGCAGAC CAACCTGGAC CGGCTGGTGG GGGATGTGCG CGAGGCCGTG TGCACGTCTG GTTGGACCTG GCCGACCACC CCCTACACGC GCTCCGGCAC						
TGCACGTCTG GTTGGACCTG GCCGACCACC CCCTACACGC GCTCCGGCAC	13201	ACGTGCAGAC	CAACCTGGAC	CGGCTGGTGG	GGGATGTGCG	CGAGGCCGTG
		TGCACGTCTG	GTTGGACCTG	GCCGACCACC	CCCTACACGC	GCTCCGGCAC

Figure 26 N

13301		TTCCTGAGTA AAGGACTCAT		CGGGGACAGG GCCCCTGTCC
13351		CAACTTTGTG GTTGAAACAC		
13401		AGGTGTACCA TCCACATGGT		
13451		CTGCAGACCG GACGTCTGGC		•
13501		GGGGGTGCGG CCCCCACGCC		
13551		CGCCCAACTC GCGGGTTGAG		
13601	••••	GGCAGCGTGT CCGTCGCACA	 	
13651		CGAGGCCATA GCTCCGGTAT		
13701		CAAGTGTCAG GTTCACAGTC		
13751		ACCCTAAACT TGGGATTTGA		
13801		CAGTTTAAAC GTCAAATTTG		
13851		TGAGCCTTAA ACTCGGAATT	 	
13901		ATGACCGCGC TACTGGCGCG		
13951	ACCGGCCGTT TGGCCGGCAA	TATCAACCGC ATAGTTGGCG		
14001	GTGAACCCCG CACTTGGGGC	AGTATTTCAC TCATAAAGTG		
14051	GCCCCCTGGT CGGGGGACCA	TTCTACACCG AAGATGTGGC		
14101	GATTCCTCTG CTAAGGAGAC	GGACGACATA CCTGCTGTAT	 	
14151	ACCCTGCTAG TGGGACGATC	AGTTGCAACA TCAACGTTGT	 •	•

7, gure 260

14251				GCTTGATAGG CGAACTATCC	
14301				GGCGAGGAGG CCGCTCCTCC	
14351				AAACCTGCCT TTTGGACGGA	
14401				AGATGAGTAG TCTACTCATC	
14451				CCGCGCCCGC	
14501				GTGGGAGGAC CACCCTCCTG	
14551				GGAGTGGCAA CCTCACCGTT	
14601				TAAAAAAAA ATTTTTTTT	
14651				CACCGAGCGT GTGGCTCGCA	
14701				TGTATGAGGA ACATACTCCT	
14751				CCAGTGGCGG GGTCACCGCC	
14801				GTTTGTGCCT CAAACACGGA	
14851	TGCGGCCTAC ACGCCGGATG	CGGGGGGAGA GCCCCCTCT	AACAGCATCC TTGTCGTAGG	GTTACTCTGA CAATGAGACT	GTTGGCACCC CAACCGTGGG
14901	CTATTCGACA GATAAGCTGT	CCACCCGTGT GGTGGGCACA	GTACCTGGTG CATGGACCAC	GACAACAAGT CTGTTGTŤCA	CAACGGATGT GTTGCCTACA
14951	GGCATCCCTG CCGTAGGGAC	AACTACCAGA TTGATGGTCT	ACGACCACAG TGCTGGTGTC	CAACTTTCTG GTTGAAAGAC	ACCACGGTCA TGGTGCCAGT
15001	TTCAAAACAA AAGTTTTGTT	TGACTACAGC ACTGATGTCG	CCGGGGGAGG GGCCCCTCC	CAAGCACACA GTTCGTGTGT	GACCATCAAT CTGGTAGTTA
15051	CTTGACGACC GAACTGCTGG	GGTCGCACTG CCAGCGTGAC	GGGCGCGAC CCCGCCGCTG	CTGAAAACCA GACTTTTGGT	TCCTGCATAC AGGACGTATG
15101	CAACATGCCA GTTGTACGGT	AATGTGAACG TTACACTTGC	AGTTCATGTT TCAAGTACAA	TACCAATAAG ATGGTTATTC	TTTAAGGCGC AAATTCCGCG



WO 02/022080 PCT/US01/28861 15151 GGGTGATGGT GTCGCGCTTG CCTACTAAGG ACAATCAGGT GGAGCTGAAA

15151	GGGTGATGGT	GTCGCGCTTG	CCTACTAAGG	ACAATCAGGT	GGAGCTGAAA
	CCCACTACCA	CAGCGCGAAC	GGATGATTCC	TGTTAGTCCA	CCTCGACTTT
•					
15201	TACGAGTGGG	TGGAGTTCAC	GCTGCCCGAG	GGCAACTACT	CCGAGACCAT
	ATGCTCACCC	ACCTCAAGTG	CGACGGGCTC	CCGTTGATGA	GGCTCTGGTA
15251	GACCATAGAC	CTTATGAACA	ACGCGATCGT	GGAGCACTAC	TTGAAAGTGG
	CTGGTATCTG	GAATACTTGT	TGCGCTAGCA	CCTCGTGATG	AACTTTCACC
15301	GCAGACAGAA	CGGGGTTCTG	GAAAGCGACA	TCGGGGTAAA	GTTTGACACC
	CGTCTGTCTT	GCCCCAAGAC	CTTTCGCTGT	AGCCCCATTT	CAAACTGTGG
15351	CGCAACTTCA	GACTGGGGTT	TGACCCCGTC	ACTGGTCTTG	TCATGCCTGG
	GCGTTGAAGT	CTGACCCCAA	ACTGGGGCAG	TGACCAGAAC	AGTACGGACC
15401	GGTATATACA	AACGAAGCCT	TCCATCCAGA	CATCATTTTG	CTGCCAGGAT
		TTGCTTCGGA			
15451	GCGGGGTGGA	CTTCACCCAC	AGCCGCCTGA	GCAACTTGTT	GGGCATCCGC
		GAAGTGGGTG			
				•	
15501	AAGCGGCAAC	CCTTCCAGGA	GGGCTTTAGG	ATCACCTACG	ATGATCTGGA
		GGAAGGTCCT	CCCGAAATCC	TAGTGGATGC	TACTAGACCT
15551	GGGTGGTAAC	ATTCCCGCAC	TGTTGGATGT	GGACGCCTAC	CAGGCGAGCT
15551		TAAGGCCGTG			
15601	TGAAAGATGA	CACCGAACAG	GGCGGGGGTG	GCGCAGGCGG	CAGCAACAGC
23002		GTGGCTTGTC			
	Actitetaçı	0100011010			010011010
15651	AGTGGCAGCG	GCGCGGAAGA	GAACTCCAAC	GCGGCAGCCG	CGGCAATGCA
15051		CGCGCCTTCT			
	10		011000110		
15701	GCCGGTGGAG	GACATGAACG	ATCATGCCAT	TCGCGGCGAC	ACCTTTGCCA
20.02		CTGTACTTGC			
15751	CACGGGCTGA	GGAGAAGCGC	GCTGAGGCCG	AAGCAGCGGC	CGAAGCTGCC
20.02		CCTCTTCGCG			
	01000001				
15801	GCCCCGCTG	CGCAACCCGA	GGTCGAGAAG	CCTCAGAAGA	AACCGGTGAT
13001		GCGTTGGGCT			
	CGGGGGGGAC	0001100001		000101101	
15851	CAAACCCCTG	ACAGAGGACA	GCAAGAAACG	CAGTTACAAC	CTAATAAGCA
13031		TGTCTCCTGT			
	GIIIGGGGAC	101010101		010.2110110	
15901	ATGACAGCAC	CTTCACCCAG	TACCGCAGCT	GGTACCTTGC	ATACAACTAC
13301		GAAGTGGGTC			
	INCIGICGIG	GANG16661C	71.0000100.		
15051	GGCGACCCTC	ል ርልርርርልልም	ררפריזר <i>≱</i> יזרבים	ACCCTGCTTT	GCACTCCTGA
10201		TCTGGCCTTA			
	CCGC1GGGAG	1C1GGCC11M	GGCGAGIACC	. Judiconin	
16001	CGTAACCTGC	CCCTCCCACC	ACCIPCITA CIPC	התיבייים ברר א	GACATGATGC
TOOUT	GCATTGGACG				
	GCA11GGACG	CCGMGCCTCG	ICCAGAIGAC	CHOCHNOO!	CIGINCINCO
16051	AAGACCCCGT	CXCCDDCCCC		ACAMCACCAA	<u>ር</u> ተተጥጥር ርርርጥር
TOOT		CTGGAAGGCG			
	TTCTGGGGCA	CIGGAAGGCG	WORLDCACE	TCTAGTCGTT	GAAAGGCCAC

Figure 26 Q

16151				TACCTCTCTG ATGGAGAGAC	
16201				CCCCCCCCC	
16251				CTCACAGATC GAGTGTCTAG	
16301	TGGCGACGCG	TTGTCGTAGC	CTCCTCAGGT	GCGAGTGACC CGCTCACTGG	TAATGACTGC
16351	GGTCTGCGGC	GTGGACGGGG	ATGCAAATGT	AGGCCCTGGG TCCGGGACCC	GTATCAGAGC
16401	GGCGCGCAGG	ATAGCTCGGC	GTGAAAAACT	GCAAGCATGT CGTTCGTACA	GGTAGGAATA
16451	TAGCGGGTCG	TTATTGTGTC	CGACCCCGGA	GCGCTTCCCA CGCGAAGGGT	TCGTTCTACA
16501	AACCGCCCCG	GTTCTTCGCG	AGGCTGGTTG	ACCCAGTGCG TGGGTCACGC	GCACGCGCCC
16551	GTGATGGCGC	GCGGGACCCC	GCGCGTGTTT	CGCGGCCGCA GCGCCGGCGT	GACCCGCGTG
16601	GTGGCAGCTA	CTGCGGTAGC	TGCGCCACCA	CCTCCTCCGC	
16651	CGCCCACGCC	GCCACCAGTG CGGTGGTCAC	TCCACAGTGG AGGTGTCACC	ACGCGGCCAT TGCGCCGGTA	TCAGACCGTG AGTCTGGCAC
16701	CACGCGCCTC	GGGCCGCGAT	ACGATTTTAC	TTCTCTGCCG	GGAGGCGCGT CCTCCGCGCA
16751	TCGTGCAGCG	GTGGCGGCGG	CTGGGCCGTG	ACGGCGGGTT	
	GCCGGGACGA	ATTGGCGCGT	GCAGCGTGGC	CGGCTGCCCG	GGCCATGCGG CCGGTACGCC
	CGGCGAGCTT	CCGACCGGCG	CCCATAACAG	TGACACGGGG	CCAGGTCCAG GGTCCAGGTC
	CGCTGCTCGC	CGGCGGCGTC	GTCGGCGCCG	GTAATCACGA	ATGACTCAGG TACTGAGTCC
	CAGCGTCCCC	GTTGCACATA	ACCCACGCGC	TGAGCCAATC	CGGCCTGCGC GCCGGACGCG
17001	GTGCCCGTGC CACGGGCACG	GCACCCGCCC CGTGGGCGGG	CCCGCGCAAC GGGCGCGTTG	TAGATTGCAA ATCTAACGTT	GAAAAAACTA CTTTTTTGAT

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17101			AAAGAAGAGA TTTCTTCTCT	
17151			GAAGGAAGAG CTTCCTTCTC	
17201			AAAAGAAAGA TTTTCTTTCT	
17251			GCTACCGCGC CGATGGCGCG	
17301		-	TGTTTTGCGA ACAAAACGCT	
17351			CCCGCACCTA GGGCGTGGAT	
17401			CTTGAGCAGG GAACTCGTCC	
17451			TAAGGACATG ATTCCTGTAC	
17501			TAAAGCCCGT ATTTCGGGCA	
17551			GAAAAGCGCG CTTTTCGCGC	
17601			GCTGATGGTA CGACTACCAT	
17651			CCGTGGAACC GGCACCTTGG	
17701	GCGTGCGGCC CGCACGCCGG			GCAGACCGTG CGTCTGGCAC
177 <u>5</u> 1	GACGTTCAGA CTGCAAGTCT			CCGCCACAGA GGCGGTGTCT
17801	GGGCATGGAG CCCGTACCTC			GCGGATGCCG CGCCTACGGC
17851	CGGTGCAGGC GCCACGTCCG			GGAGGTGCAA CCTCCACGTT
17901	ACGGACCCGT TGCCTGGGCA			CGCGCCGTTC GCGCGGCAAG
17951	GAGGAAGTAC CTCCTTCATG			GCCCTACATC CGGGATGTAG

Figure 265

18051		CTACCCGACG GATGGGCTGC			
18101		CAGCCCGTGC GTCGGGCACG			
18151		CAGGACCCTG GTCCTGGGAC			
18201		AGCCGGTCTT TCGGCCAGAA			
18251		TTCCCGGTGC AAGGGCCACG			
18301		CGGCCACGGC GCCGGTGCCG			
18351	GCCGCCGCCG	GCGCGTCGCA CGCGCAGCGT	GGCAGCGTAC	GCGCCGCCAT	AGGACGGGGA
18401		CTGATCGCCG GACTAGCGGC			
18451		GCAGGCGCAG CGTCCGCGTC			
18501		ATAAAAAGTC TATTTTTCAG			
18551		AATGGAAGAC TTACCTTCTG			
18601		CGTTCATGGG GCAAGTACCC			
18651	GAGCGGTGGC CTCGCCACCG	GCCTTCAGCT CGGAAGTCGA	GGGGCTCGCT CCCCGAGCGA	GTGGAGCGGC CACCTCGCCG	TTAAAAATTAAATTTAAATTTTAAA
18701	TCGGTTCCAC AGCCAAGGTG	CGTTAAGAAC GCAATTCTTG	TATGGCAGCA ATACCGTCGT	AGGCCTGGAA TCCGGACCTT	CAGCAGCACA GTCGTCGTGT
18751	GGCCAGATGC CCGGTCTACG	TGAGGGATAA ACTCCCTATT	GTTGAAAGAG CAACTTTCTC	CAAAATTTCC GTTTTAAAGG	AACAAAAGGT TTGTTTTCCA
18801	GGTAGATGGC CCATCTACCG	CTGGCCTCTG GACCGGAGAC	GCATTAGCGG CGTAATCGCC	GGTGGTGGAC CCACCACCTG	CTGGCCAACC GACCGGTTGG
18851	AGGCAGTGCA TCCGTCACGT	AAATAAGATT TTTATTCTAA	AACAGTAAGC TTGTCATTCG	TTGATCCCCG AACTAGGGGC	CCCTCCCGTA GGGAGGGCAT
18901	GAGGAGCCTC CTCCTCGGAG	CACCGGCCGT GTGGCCGGCA	GGAGACAGTG CCTCTGTCAC	TCTCCAGAGG AGAGGTCTCC	GGCGTGGCGA CCGCACCGCT

Figure 26T

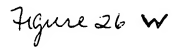
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19151		CGTTGTTGTA GCAACAACAT		
19201		GTCCGCGATC CAGGCGCTAG		
19251		AACAGCATCG TTGTCGTAGC		
19301	±	CTGATAGCTA GACTATCGAT		
19351		CAGAGGAGCT GTCTCCTCGA		
19401		TTCGATGATG AAGCTACTAC		
19451		CGGAGTACCT GCCTCATGGA		
19501		TACTTCAGCC ATGAAGTCGG		
19551		CGACGTGACC GCTGCACTGG		
19601		TGGACCGTGA ACCTGGCACT		AGGCGCGGTT TCCGCGCCAA
19651	CACCCTAGCT GTGGGATCGA			TCCACGTACT AGGTGCATGA
19701	TTGACATCCG AACTGTAGGC			GCCCTACTCT CGGGATGAGA
19751	GGCACTGCCT CCGTGACGGA	ACAACGCCCT TGTTGCGGGA		
19801	ATGGGATGAA TACCCTACTT	GCTGCTACTG CGACGATGAC		
19851	ATGACAACGA TACTGTTGCT	AGACGAAGTA TCTGCTTCAT		

Figure 26 U

19951			AAACACCTAA TTTGTGGATT		
20001			GAATCTCAGT CTTAGAGTCA		
20051			AAAAAAGACT TTTTTTCTGA		
20101			CAAATGAAAA GTTTACTTTT		
20151			CTAGAAAGTC GATCTTTCAG		
20201	TCAACTACTG AGTTGATGAC	AGGCAGCCGC TCCGTCGGCG	AGGCAATGGT TCCGTTACCA	GATAACTTGA CTATTGAACT	CTCCTAAAGT GAGGATTTCA
20251	GGTATTGTAC CCATAACATG	AGTGAAGATG TCACTTCTAC	TAGATATAGA ATCTATATCT	AACCCCAGAC TTGGGGTCTG	ACTCATATTT TGAGTATAAA
20301	GAATGTACGG	GTGATAATTC	GAAGGTAACT CTTCCATTGA	GTGCTCTTGA	TTACCCGGTT
20351	GTTAGATACG	GGTTGTCCGG	TAATTACATT ATTAATGTAA	CGAAAATCCC	TGTTAAAATA
20401	ACCAGATTAC	ATAATGTTGT	GCACGGGTAA CGTGCCCATT	ATACCCACAA	GACCGCCCGG
20451	TTCGTAGCGT	CAACTTACGA	GTTGTAGATT CAACATCTAA	ACGTTCTGTC	TTTGTGTCTC
20501	GAAAGTATGG	TCGAAAACGA	TGATTCCATT ACTAAGGTAA	CCACTATCTT	GGTCCATGAA
20551	AAGATACACC	TTAGTCCGAC	AACTGTCGAT	ACTAGGTCTA	
	AACTTTTAGT	ACCTTGACTT	CTACTTGAAG	GTTTAATGAC	CTTTCCACTG GAAAGGTGAC
	CCTCCACACT	AATTATGTCT	CTGAGAATGG	TTCCATTTTG	CTAAAACAGG GATTTTGTCC
20701	TCAGGAAAAT AGTCCTTTTA	GGATGGGAAA	AAGATGCTAC TTCTACGATG	AGAATTTTCA TCTTAAAAGT	GATAAAAATG CTATTTTTAC
20751					AAATGCCAAC TTTACGGTTG
20801	CTGTGGAGAA GACACCTCTT	ATTTCCTGTA TAAAGGACAT	CTCCAACATA GAGGTTGTAT	GCGCTGTATT CGCGACATAA	TGCCCGACAA ACGGGCTGTT

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20901			CCGGGCTAGT GGCCCGATCA	GGACTGCTAC CCTGACGATG
20951		-	TATATGGACA ATATACCTGT	- :
21001			CTACCGCTCA GATGGCGAGT	
21051			AGGTGCCTCA TCCACGGAGT	
21101			TCATACACCT AGTATGTGGA	
21151			GAGCTCCCTA CTCGAGGGAT	
21201	-	 	ATAGCATTTG TATCGTAAAC	
21251		 	TCCACGCTTG AGGTGCGAAC	
21301		 	CGACTATCTC GCTGATAGAG	
21351			CCAACGTGCC GGTTGCACGG	
21401		 	TGGGCCTTCA ACCCGGAAGT	
21451			CTACGACCCT GATGCTGGGA	
21501			CCTTTTACCT GGAAAATGGA	
21551	TTTAAGAAGG AAATTCTTCC		TCTGTCAGCT AGACAGTCGA	
21601	TGACCGCCTG ACTGGCGGAC		AATTAAGCGC TTAATTCGCG	
21651	GGGAGGGTTA CCCTCCCAAT		TGACCAAAGA ACTGGTTTCT	
21701	GTACAAATGC CATGTTTACG		TACCAGGGCT ATGGTCCCGA	
21751	AGAGAGCTAC TCTCTCGATG		CTTTAGAAAC GAAATCTTTG	



21851		ACCAACACAA TGGTTGTGTT			
21901		GAAGGACAGG CTTCCTGTCC			
21951		CGCAGTTGAC GCGTCAACTG			
22001		GGCGCATCCC CCGCGTAGGG			
22051		CTGGGCCAAA GACCCGGTTT			
22101		TTTTGAGGTG AAAACTCCAC			
22151		AAGTCTTTGA TTCAGAAACT			
22201		ACCGTGTACC TGGCACATGG			
22251		AAGCAAGCAA TTCGTTCGTT			
22301		ACTGAAAGCC TGACTTTCGG			
22351	TTTTTGGGCA AAAAACCCGT	CCTATGACAA GGATACTGTT	GCGCTTTCCA CGCGAAAGGT	GGCTTTGTTT CCGAAACAAA	CTCCACACAA GAGGTGTGTT
22401	CGAGCGGACG	GCCATAGTCA CGGTATCAGT	TATGCCGGCC	AGCGCTCTGA	CCCCCGCATG
22451	ACTGGATGGC TGACCTACCG	CTTTGCCTGĠ GAAACGGACC	AACCCGCACT TTGGGCGTGA	CAAAAACATG GTTTTTGTAC	CTACCTCTTT GATGGAGAAA
22501					ACCAGTTTGA TGGTCAAACT
22551	GTACGAGTCA CATGCTCAGT	CTCCTGCGCC GAGGACGCGG	GTAGCGCCAT CATCGCGGTA	TGCTTCTTCC ACGAAGAAGG	CCCGACCGCT GGGCTGGCGA
22601	GTATAACGCT CATATTGCGA	GGAAAAGTCC CCTTTTCAGG	ACCCAAAGCG TGGGTTTCGC	TACAGGGGCC ATGTCCCCGG	CAACTCGGCC GTTGAGCCGG
	CGGACACCTG	ATAAGACGAC	GTACAAAGAG	GTGCGGAAAC	CCAACTGGCC GGTTGACCGG
22701	CCAAACTCCC GGTTTGAGGG	ATGGATCACA TACCTAGTGT	ACCCCACCAT TGGGGTGGTA	GAACCTTATT CTTGGAATAA	ACCGGGGTAC TGGCCCCATG

Figure 26 X

22801	GTCCTTGTCG	AGATGTCGAA	GGACCTCGCG	CACTOGOCOT GTGAGCGGGA	TGAAGGCGTC
22851	GGTGTCACGC	GTCTAATCCT	CGCGGTGAAG	TTTTTGTCAC AAAAACAGTG	AACTTTTTGT
22901				ATAAAGGCAA TATTTCCGTT	
22951				ACCCTTGCCG TGGGAACGGC	
23001	AATTTTTAGT	TTCCCCAAGA	CGGCGCGTAG	GCTATGCGCC CGATACGCGG	TGACCGTCCC
23051	TGTGCAACGC	TATGACCACA	AATCACGAGG	ACTTAAACTC TGAATTTGAG	TCCGTGTTGG
23101	TAGGCGCCGT	CGAGCCACTT	CAAAAGTGAG	CACAGGCTGC GTGTCCGACG	CGTGGTAGTG
23151	GTTGCGCAAA	TCGTCCAGCC	CGCGGCTATA	CTTGAAGTCG GAACTTCAGC	GTCAACCCCG
23201	GAGGCGGGAC	GCGCGCGCTC	AACGCTATGT	CAGGGTTGCA GTCCCAACGT	CGTGACCTTG
23251	TGATAGTCGC	GGCCCACCAC	GTGCGACCGG	AGCACGCTCT TCGTGCGAGA	ACAGCCTCTA
23301	GTCTAGGCGC	AGGTCCAGGA	GGCGCAACGA	CAGGGCGAAC GTCCCGCTTG	CCTCAGTTGA
23351	AACCATCGAC	GGAAGGGTTT	TTCCCGCGCA	GCCCAGGCTT CGGGTCCGAA	ACTCAACGTG
23401	AGCGTGGCAT	CACCGTAGTT	TTCCACTGGC		CCCGCAATCC
	TATGTCGCGG	ACGTATTTTC	GGAACTAGAC	GAATTTTCGG	ACCTGAGCCT TGGACTCGGA
•	AACGCGGAAG	TCTCTTCTTG	TACGGCGTTC	TGAACGGCCT	AAACTGATTG TTTGACTAAC
23551	GCCGGACAGG CGGCCTGTCC	CCGCGTCGTG	CACGCAGCAC GTGCGTCGTG	CTTGCGTCGG GAACGCAGCC	TGTTGGAGAT ACAACCTCTA
23601	CTGCACCACA GACGTGGTGT	TTTCGGCCCC AAAGCCGGGG	ACCGGTTCTT TGGCCAAGAA	CACGATCTTG GTGCTAGAAC	GCCTTGCTAG CGGAACGATC
23651	ACTGCTCCTT TGACGAGGAA	CAGCGCGCGC GTCGCGCGCGC	TGCCCGTTTT ACGGGCAAAA	CGCTCGTCAC GCGAGCAGTG	ATCCATTTCA TAGGTAAAGT

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23701	ATCACGTGCT TAGTGCACGA			GGCACATCTG	
23751	GCCTTCGATC	TCAGCGCAGC	GGTGCAGCCA	CAACGCGCAG	CCCGTGGGCT
	CGGAAGCTAG	AGTCGCGTCG	CCACGTCGGT	GTTGCGCGTC	GGGCACCCGA
23801	CGTGATGCTT	GTAGGTCACC	TCTGCAAACG	ACTGCAGGTA	CGCCTGCAGG
	GCACTACGAA	CATCCAGTGG	AGACGTTTGC	TGACGTCCAT	GCGGACGTCC
. 23851	AATCGCCCCA	TCATCGTCAC	AAAGGTCTTG	TTGCTGGTGA	AGGTCAGCTG
	TTAGCGGGGT	AGTAGCAGTG	TTTCCAGAAC	AACGACCACT	TCCAGTCGAC
23901	CAACCCGCGG	TGCTCCTCGT	TCAGCCAGGT	CTTGCATACG	GCCGCCAGAG
	GTTGGGCGCC	ACGAGGAGCA	AGTCGGTCCA	GAACGTATGC	CGGCGGTCTC
23951	CTTCCACTTG	GTCAGGCAGT	AGTTTGAAGT	TCGCCTTTAG	ATCGTTATCC
	GAAGGTGAAC	CAGTCCGTCA	TCAAACTTCA	AGCGGAAATC	TAGCAATAGG
24001	ACGTGGTACT				
				CGGAGGTACG	
24051	CGCAGACACG			=	
				GTAGTGGCAT	
24101				GCGTCCGCAT	
	•			CGCAGGCGTA	
24151	_			GTGCGCTTAC	
				CACGCGAATG	
24201				ACCCACCATT	
24251	TACGAACTAA			TGGGTGGTAA	
24251				AATGGAGACC	
24301	CGCTCGGGCT				
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24351	CAAATCCGCC	GCCGAGGTCG	ATGGCCGCGG	GCTGGGTGTG	CGCGGCACCA
				CGACCCACAC	
24401	GCGCGTCTTG				
				GCCTGAGCTA	
24451	ATCCGCTTTT				
				CCGCCGCTGC	
24501	CGACÁCGTCC				
				GCGGCGTGGC	
24551	CGGGGGTGGT				
				CTGACCGGTA	
24601	TATAGGCAGA				
	ATATCCGTCT	TTTTCTAGTA	CCTCAGTCAG	CTCTTCTTCC	TGTCGGATTG

Figure 262 80/144

24701					GGAAGTGATT CCTTCACTAA
24751		ACCCAGGTTT TGGGTCCAAA			ACCGCTCAGT TGGCGAGTCA
24801		GATAAAAGC CTATTTTTCG			GCAAACGAGG CGTTTGCTCC
24851	AACAAGTCGG TTGTTCAGCC	GCGGGGGGAC CGCCCCCTG			
24901		TGTTGAAGCA ACAACTTCGT			
24951		GAGCGCAGCG CTCGCGTCGC			
25001		ACGCCACCTA TGCGGTGGAT			
25051		CATGCGAGCC GTACGCTCGG			
25101		GAGGTGCTTG CTCCACGAAC			
25151		ATCCTGCCGT TAGGACGGCA			
25201		AGGGCGCTGT TCCCGCGACA			
25251		TTTGAGGGTC AAACTCCCAG			
25301	CTCTGCAACA GAGACGTTGT				AGTGTTGGTG TCACAACCAC
25351	GAACTCGAGG CTTGAGCTCC	GTGACAACGC CACTGTTGCG	GCGCCTAGCC CGCGGATCGG	GTACTAAAAC CATGATTTTG	GCAGCATCGA CGTCGTAGCT
25401	GGTCACCCAC CCAGTGGGTG	TTTGCCTACC AAACGGATGG			
25,451	GCACAGTCAT CGTGTCAGTA				CCTGGAGAGG GGACCTCTCC
25501	GATGCAAATT CTACGTTTAA	TGCAAGAACA ACGTTCTTGT	AACAGAGGAG TTGTCTCCTC	GGCCTACCCG CCGGATGGGC	CAGTTGGCGA GTCAACCGCT
25551	CGAGCAGCTA GCTCGTCGAT	GCGCGCTGGC CGCGCGACCG	TTCAAACGCG AAGTTTGCGC	CGAGCCTGCC GCTCGGACGG	GACTTGGAGG CTGAACCTCC

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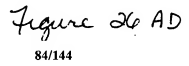
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25751			CCTACCTTGG GGATGGAACC	
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25851			CTTATTTCTA GAATAAAGAT	
25901			GCTTGGAGGA CGAACCTCCT	
25951	AAGGAGCTGC TTCCTCGACG		TTGAAGGACC AACTTCCTGG	
26001			GGCGGACATC CCGCCTGTAG	
26051			TGCCAGACTT ACGGTCTGAA	
26101			CTAGAGCGCT GATCTCGCGA	
26151			CTTTGTGCCC GAAACACGGG	
26201			GCTACCTTCT CGATGGAAGA	
26251	AACTACCTTG TTGATGGAAC	 	-	GCGGTGACGG CGCCACTGCC
26301	TCTACTGGAG AGATGACCTC			CACCGCTCCC GTGGCGAGGG
26351	TGGTTTGCAA ACCAAACGTT			CGGTACCTTT GCCATGGAAA
26401	GAGCTGCAGG CTCGACGTCC			CGGGGTTGAA GCCCCAACTT
26451	ACTCACTCCG TGAGTGAGGC			TTTGTACCTG AAACATGGAC
26501	AGGACTACCA TCCTGATGGT			ATCCCGCCCG TAGGGCGGGC

Figure 26 AB

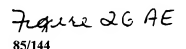
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26601	CCAATTGCAA	GCCATCAACA	AAGCCCGCCA	AGAGTTTCTG	CTACGAAAGG
	GGTTAACGTT	CGGTAGTTGT	TTCGGGCGGT	TCTCAAAGAC	GATGCTTTCC
26651	GACGGGGGT				
				CGCTCCTCGA	
26701				CCGCGGGCCC	
	GGGGGCGGCG	GCGTCGGGAT	AGICGICGIC	GGCGCCCGGG	AACGAAGGGI
26751	GGATGGCACC	CAAAAAGAAG	CTGCAGCTGC	CGCCGCCACC	CACGGACGAG
	CCTACCGTGG	GTTTTTCTTC	GACGTCGACG	GCGCCGTGG	GTGCCTGCTC
26801	GAGGAATACT				
				CAAAACCTGC	
26851	GGACATGATG			CGAGGAAGCT	
26901	AAGAGGTGTC				
	TTCTCCACAG	TCTGCTTTGT	GGCAGTGGGA	GCCAGCGTAA	GGGGAGCGGC
26951	GCGCCCCAGA				
				TACCGATGTT	
27001	TCAGGCGCCG				
				TGGGTTGGCA	
27051	CCACTGGAAC				
				TCGGCGGCGG	
27101	GAGCAACAAC			TGGCGCGGGC	
	0.00				
27151	CATAGTTGCT			CAACATCTCC	
27201	GCTTTCTTCT				The state of the s
					GTAGGACGTA
· 27251	TACTACCGTC				
				TGGCCGCCGT	
27301	CAGCAGCGGC				
	GTCGTCGCCG	GTGTGTCTTC	GTTTCCGCTG	GCCTATCGTT	CTGAGACTGT
27351	AAGCCCAAGA				
	TTCGGGTTCT	TTAGGTGTCG	CCGCCGTCGT	CGTCCTCCTC	CTCGCGACGC
27401	TCTGGCGCCC				
	AGACCGCGGG	TTGCTTGGGC	ATAGCTGGGC	GCTCGAATCT	TTGTCCTAAA
27451	TTCCCACTCT	GTATGCTATA	TTTCAACAGA	GCAGGGGCCA	AGAACAAGAG
	AAGGGTGAGA	CATACGATAT	AAAGTTGTCT	CGTCCCCGGT	TCTTGTTCTC

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27551			TTCGGCGCAC AAGCCGCGTG		
27601			CTGACTCTTA GACTGAGAAT		
27651			ACTACGTCAT TGATGCAGTA		
27701			CCATTATGAG GGTAATACTC		
27751			CAAATGGGAC GTTTACCCTG		
27801			CTACATGAGC GATGTACTCG		
27851			CCCACCGAAA GGGTGGCTTT		
27901			CGTAATAACC GCATTATTGG		
27951	CGACGGGACC	ACATGGTCCT	AAGTCCCGCT TTCAGGGCGA	GGGTGGTGAC	ACCATGAAGG
28001	GTCTCTGCGG	GTCCGGCTTC	TTCAGATGAC AAGTCTACTG	ATTGAGTCCC	CGCGTCGAAC
28051	GCCCGCCGAA	AGCAGTGTCC	GTGCGGTCGC CACGCCAGCG	GGCCCGTCCC	ATATTGAGTG
28101	GACTGTTAGT	CTCCCGCTCC	TATTCAGCTC ATAAGTCGAG	TTGCTGCTCA	GCCACTCGAG
		GAGGCAGGCC	TGCCCTGTAA	AGTCTAGCCG	CCGCGGCCGG
	CGAGAAGTAA	GTGCGGAGCA	GTCCGTTAGG	ATTGAGACGT	GACCTCGTCC CTGGAGCAGG
	AGACTCGGCG	CGAGACCTCC	GTAACCTTGA	GACGTTAAAT	TTGAGGAGTT AACTCCTCAA
	ACACGGTAGC	CAGATGAAAT	TGGGGAAGAG	CCCTGGAGGG	GGCCACTATC CCGGTGATAG
	GCCTAGTTAA	ATAAGGATTG	AAACTGCGCC	ATTTCCTGAG	GGCGGACGGC
28401	TACGACTGAA ATGCTGACTT	TGTTAAGTGG ACAATTCACC	AGAGGCAGAG TCTCCGTCTC	CAACTGCGCC GTTGACGCGG	TGAAACACCT ACTTTGTGGA



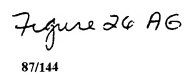
WO 02/022080 PCT/US01/28861 28451 GGTCCACTGT CGCCGCCACA AGTGCTTTGC CCGCGACTCC GGTGAGTTTT CCAGGTGACA GCGGCGTGT TCACGAAACG GGCGCTGAGG CCACTCAAAA 28501 GCTACTITGA ATTGCCCGAG GATCATATCG AGGGCCCGGC GCACGGCGTC CGATGAAACT TAACGGGCTC CTAGTATAGC TCCCGGGCCG CGTGCCGCAG 28551 CGGCTTACCG CCCAGGGAGA GCTTGCCCGT AGCCTGATTC GGGAGTTTAC GCCGAATGGC GGGTCCCTCT CGAACGGGCA TCGGACTAAG CCCTCAAATG 28601 CCAGCGCCC CTGCTAGTTG AGCGGGACAG GGGACCCTGT GTTCTCACTG GGTCGCGGGG GACGATCAAC TCGCCCTGTC CCCTGGGACA CAAGAGTGAC 28651 TGATTTGCAA CTGTCCTAAC CCTGGATTAC ATCAAGATCT TTGTTGCCAT ACTAAACGTT GACAGGATTG GGACCTAATG TAGTTCTAGA AACAACGGTA 28701 CTCTGTGCTG AGTATAATAA ATACAGAAAT TAAAATATAC TGGGGCTCCT GAGACACGAC TCATATTATT TATGTCTTTA ATTTTATATG ACCCCGAGGA 28751 ATCGCCATCC TGTAAACGCC ACCGTCTTCA CCCGCCCAAG CAAACCAAGG TAGCGGTAGG ACATTTGCGG TGGCAGAAGT GGGCGGGTTC GTTTGGTTCC 28801 CGAACCTTAC CTGGTACTTT TAACATCTCT CCCTCTGTGA TTTACAACAG GCTTGGAATG GACCATGAAA ATTGTAGAGA GGGAGACACT AAATGTTGTC 28851 TTTCAACCCA GACGGAGTGA GTCTACGAGA GAACCTCTCC GAGCTCAGCT AAAGTTGGGT CTGCCTCACT CAGATGCTCT CTTGGAGAGG CTCGAGTCGA 28901 ACTCCATCAG AAAAAACACC ACCCTCCTTA CCTGCCGGGA ACGTACGAGT TGAGGTAGTC TTTTTTGTGG TGGGAGGAAT GGACGGCCCT TGCATGCTCA 28951 GCGTCACCGG CCGCTGCACC ACACCTACCG CCTGACCGTA AACCAGACTT CGCAGTGGCC GGCGACGTGG TGTGGATGGC GGACTGGCAT TTGGTCTGAA 29001 TTTCCGGACA GACCTCAATA ACTCTGTTTA CCAGAACAGG AGGTGAGCTT AAAGGCCTGT CTGGAGTTAT TGAGACAAAT GGTCTTGTCC TCCACTCGAA 29051 AGAAAACCCT TAGGGTATTA GGCCAAAGGC GCAGCTACTG TGGGGTTTAT TCTTTTGGGA ATCCCATAAT CCGGTTTCCG CGTCGATGAC ACCCCAAATA 29101 GAACAATTCA AGCAACTCTA CGGGCTATTC TAATTCAGGT TTCTCTAGAA CTTGTTAAGT TCGTTGAGAT GCCCGATAAG ATTAAGTCCA AAGAGATCTT 29151 TCGGGGTTGG GGTTATTCTC TGTCTTGTGA TTCTCTTTAT TCTTATACTA AGCCCCAACC CCAATAAGAG ACAGAACACT AAGAGAAATA AGAATATGAT 29201 ACGCTTCTCT GCCTAAGGCT CGCCGCCTGC TGTGTGCACA TTTGCATTTA TGCGAAGAGA CGGATTCCGA GCGGCGGACG ACACACGTGT AAACGTAAAT 29251 TTGTCAGCTT TTTAAACGCT GGGGTCGCCA CCCAAGATGA TTAGGTACAT AACAGTCGAA AAATTTGCGA CCCCAGCGGT GGGTTCTACT AATCCATGTA 29301 AATCCTAGGT TTACTCACCC TTGCGTCAGC CCACGGTACC ACCCAAAAGG TTAGGATCCA AATGAGTGGG AACGCAGTCG GGTGCCATGG TGGGTTTTCC 29351 TGGATTTTAA GGAGCCAGCC TGTAATGTTA CATTCGCAGC TGAAGCTAAT ACCTAAAATT CCTCGGTCGG ACATTACAAT GTAAGCGTCG ACTTCGATTA



29451		AACAAAATTG TTGTTTTAAC			
29501		TACAGAGTAT ATGTCTCATA			
29551		TGTATACTTT ACATATGAAA			
29601		AAACAGTATA TTTGTCATAT			
29651	-	TTTCTGCTGC AAAGACGACG			
29701		TACTCTATAT ATGAGATATA			
29751		ATGCCTTAAT TACGGAATTA			
29801		CTCGCTGCTT GAGCGACGAA			
29851		GATTTAAACC CTAAATTTGG			
29901	-	TGACTCTATG ACTGAGATAC			
29951	••••	CTGGATGTCA GACCTACAGT			
30001		CAGTCCAACT GTCAGGTTGA			
30051	CACAACCAAC GTGTTGGTTG	CGCCGCCGC			
		ACGGAAACAG	TTATTGACCC	TATTGAACCC	GTACACCACC
		GCGAATACAA	ACATACGGAA	TAATAATACA	CCGAGTAGAC
		GCGTTTGCGC	GGGCTGGTGG	GTAGATATCA	GGGTAGTAAC
		TTTGTTACTA	CCTTAGGTAT	CTAACCTGCC	TGACTTTGTG
30301	ATGTTCTTTT TACAAGAAAA	CTCTTACAGT GAGAATGTCA			

Figure 26 AF

30401				TCCAGCCTTC AGGTCGGAAG	ACAGTCTATT TGTCAGATAA
30451				TCTGCAGCCT AGACGTCGGA	
30501				GTCTGTGTGC CAGACACACG	GCTTTGCATA CGAAACGTAT
.30551				GACTATAGCT CTGATATCGA	
30601	-			TTTTCTGCTG AAAAGACGAC	
30651				AGCCTCAAAG TCGGAGTTTC	
30701				AGTTGCTACA TCAACGATGT	
30751				CATCTCTGTT GTAGAGACAA	
30801				CCTACCTTGA GGATGGAACT	
30851	TTGCGTTATC	TACGGTACTT	GGTGGGTTGA	TTCCCCGCGC AAGGGGCGCG	GGCGATACGA
30901				TGTCCCAGCC ACAGGGTCGG	
30951				GCTACTTTAA CGATGAAATT	
	•	CTGTGGGATC	TAGATCTTTÀ	CCTGCCTTAA	TAATGTCTCG
		TCTTTCTGCG	TCCCGTCGCC	GGCTCGTTGT	CGCGTACTTA
		TTCTGTACCA	ATTGAACGTG	GTCACGTTTT	CCCCATAGAA
		TTCGTCCGGT	TTCAGTGGAT	GCTGTCATTA	TGGTGGCCTG
		GATGTTCAAC	GGTTGGTTCG	CAGTCTTTAA	CCACCAGTAC
31251	GTGGGAGAAA CACCCTCTTT				AAACCGAAGG TTTGGCTTCC



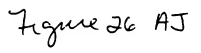
31351					AAAAAAAAA TTTTTTTTAT
31401					CTGTCCAGTT GACAGGTCAA
31451					TTGCAGCTTC AACGTCGAAG
31501			CCACAATCTA GGTGTTAGAT		CAGTTTCCTC GTCAAAGGAG
31551			CCACTATCTT GGTGATAGAA		
31601			ACCTTCAACC TGGAAGTTGG		
31651			GCCTTTTCTT CGGAAAAGAA		=
31701			CCCCTGGGGT GGGGACCCCA		
31751			GGCATGCTTG CCGTACGAAC		
31801			CAACCTTACC GTTGGAATGG		
31851			CCAAGTCAAA GGTTCAGTTT		
31901			GAAGCCCTAA CTTCGGGATT		
	CTAATGGTCG GATTACCAGC	•			
32001	CGTGCACGAC GCACGTGCTG				
32051	CAGAAGGAAA GTCTTCCTTT				CACCACCGAT GTGGTGGCTA
32101	AGCAGTACCC TCGTCATGGG		· -	-	
32151	TAGCTTGGGC ATCGAACCCG				
	TAGGACTAAA ATCCTGATTT				

Figure 26 AH

32301	AACTAAAGTT TTGATTTCAA	ACTGGAGCCT TGACCTCGGA			
32351	TTAATGTAGC AATTACATCG	AGGAGGACTA TCCTCCTGAT			
32401	CTTGATGTTA GAACTACAAT	GTTATCCGTT CAATAGGCAA			
32451	AGGACAGGGC TCCTGTCCCG	CCTCTTTTTA GGAGAAAAT			
32501		CCTTTACTTG GGAAATGAAC			
32551		TAAGCACTGC ATTCGTGACG			
32601		GCAGGAGATG CGTCCTCTAC			
32651		CCTCAAAACA GGAGTTTTGT			
32701	AACAAGGCTA TTGTTCCGAT	TGGTTCCTAA ACCAAGGATT			
32751		ACAGTAGGAA TGTCATCCTT			
32801		TCCATCTCCT AGGTAGAGGA			
32851	TTTGAGTGAA	TGGTCTTAAC ACCAGAATTG	TTTTACACCG	TCAGTTTATG	AACGATGTCA
32901	TTCAGTTTTG AAGTCAAAAC	GCTGTTAAAG CGACAATTTC	GCAGTTTGGC CGTCAAACCG	TCCAATATCT AGGTTATAGA	GGAACAGTTC CCTTGTCAAG
	TTTCACGAGT	AGAATAATAT	TCTAAACTGC	TTTTACCTCA	GCTACTAAAC CGATGATTTG
	TTAAGGAAGG	ACCTGGGTCT	TATAACCTTG	AAATCTTTAC	GAGATCTTAC CTCTAGAATG
		CGGATATGTT	TGCGACAACC	TAAATACGGA	TTGGATAGTC
	GAATAGGTTT	TAGAGTGCCA	TTTTGACGGT	TTTCATTGTA	TGTCAGTCAA ACAGTCAGTT
33151	GTTTACTTAA CAAATGAATT	ACGGAGACAA TGCCTCTGTT	AACTAAACCT TTGATTTGGA	GTAACACTAA CATTGTGATT	CCATTACACT GGTAATGTGA

Figure 26 AI 89/144

33251		GGACTGGTCT CCTGACCAGA			
33301		ACACTTTTTC TGTGAAAAAG			
33351		CAACGTGTTT GTTGCACAAA			
33401		GTAGTATAGC CATCATATCG			
33451		CAAACTCACA GTTTGAGTGT			
33501		AGAGTACACA TCTCATGTGT			
33551		GGGTAACAGA CCCATTGTCT			
33601		GCCAAACGCT CGGTTTGCGA			
33651	CGAGTGAATT	GTTCATGTCG CAAGTACAGC	GACAGGTCGA	CGACTCGGTG	TCCGACGACA
33701	GGTTGAACGC	GTTGCTTAAC CAACGAATTG	CCCGCCGCTT	CCTCTTCAGG	TGCGGATGTA
33751	CCCCATCTC	TCATAATCGT AGTATTAGCA	CGTAGTCCTA	TCCCGCCACC	ACGACGTCGT
33801	CGCGCGCTTA	AAACTGCTGC TTTGACGACG	GCGGCGGCGA	GGCAGGACGT	CCTTATGTTG
		AGAGGAGTCG	CTACTAAGCG	TGGCGGGCGT	CGTATTCCGC
	GGAACAGGAG	GCCCGTGTCG	TCGCGTGGGA	CTAGAGTGAA	AAATCAGCAC TTTAGTCGTG
	TCATTGACGT	CGTGTCGTGG	TGTTATAACA	AGTTTTAGGG	ACAGTGCAAG TGTCACGTTC
	CGCGACATAG	GTTTCGAGTA	CCGCCCCTGG	TGTCTTGGGT	CGTGGCCATC GCACCGGTAG
	TATGGTGTTC	GCGTCCATCT	AATTCACCGC	TGGGGAGTAT	AACACGCTGG TTGTGCGACC
34101	ACATAAACAT TGTATTTGTA	TACCTCTTTT ATGGAGAAAA	GGCATGTTGT CCGTACAACA	AATTCACCAC TTAAGTGGTG	CTCCCGGTAC GAGGGCCATG



34201		ACCTGCCCGC TGGACGGGCG		
34251		GTGGAGAGCC CACCTCTCGG		
34301		CAATGTTGGC GTTACAACCG		TACACTTCCT ATGTGAAGGA
34351	CAGGATTACA GTCCTAATGT	AGCTCCTCCC TCGAGGAGGG		
34401		CAGCGTAAAT GTCGCATTTA		
34451		GCATTGTCAA CGTAACAGTT		
34501		GTAGCGCGGG CATCGCGCCC		
34551		AGTGCGCCGA TCACGCGGCT		
	ATGCCAAATG TACGGTTTAC			
34651		ACAAACAGAT TGTTTGTCTA		
34701		AGTTGTAGTA TCAACATCAT		
34751		GGTTCTATGT CCAAGATACA		
34801	CATCCACCAC GTAGGTGGTG			ACATTCGTTC TGTAAGCAAG
	TGCGAGTCAC ACGCTCAGTG			
	TTTTTTATTC AAAAAATAAG			
34951	TGAACGCGCT ACTTGCGCGA			CCAAAGAACA GGTTTCTTGT
35001	GATAATGGCA CTATTACCGT			AGGCAAACGG TCCGTTTGCC
35051	CCCTCACGTC GGGAGTGCAG			GTGAATCTCC CACTTAGAGG

Figure 26 AK

35151			CCGAATATTA GGCTTATAAT	
35201			CCTTCAGCCT GGAAGTCGGA	
35251			AGACCTGTAT TCTGGACATA	
35301			CGTAGGTCCC GCATCCAGGG	
35351			GACCAGCGCG CTGGTCGCGC	
35401			TGATTATGAC ACTAATACTG	
35451			TAAGCTTGTT ATTCGAACAA	
35501			ATCAGGCAAA TAGTCCGTTT	
35551	AAAAAGAAAG TTTTTCTTTC		GCAGATAAAG CGTCTATTTC	
35601			TTTCTCTCAA AAAGAGAGTT	
35651			CAAAAAAACA GTTTTTTTGT	
35701			CCTTATAAGC GGAATATTCG	
35751	CTACGGCCAT GATGCCGGTA			GTGATTAAAA CACTAATTTT
35801	AGCACCACCG TCGTGGTGGC		GGAGTCATAA CCTCAGTATT	
35851	GGTAAACACA CCATTTGTGT		CAGTGCTAAA GTCACGATTT	
35901	AATAGCCCGG TTATCGGGCC			CATTACAGCC GTAATGTCGG
35951	CCCATAGGAG GGGTATCCTC			CATAAACACC GTATTTGTGG
36001	TGAAAAACCC ACTTTTTGGG			TCCAGAACAA AGGTCTTGTT

Figure 26 AL

36101				ACACGGCACC TGTGCCGTGG	
36151	* · · · · ·			GCGAGTATAT CGCTCATATA	
36201	AAAATGACGT TTTTACTGCA			AACACCCAGA TTGTGGGTCT	
36251				AACCCACAAC TTGGGTGTTG	
36301	GCAGTGAAGG	CAAAAGGGTG	CAATGCAGTG	TTCCCATTTT AAGGGTAAAA	TTCTTTTGAT
36351	GTTAAGGGTT	GTGTATGTTC	AATGAGGCGG	CTAAAACCTA GATTTTGGAT	GCAGTGGGCG
36401				ACTCCACCCC TGAGGTGGGG	GAGTAATAGT
					PacI
36451	TATTGGCTTC				
				ТААСТАСТАС	
36501				CTTCCCCATT GAAGGGGTAA	
36551				TGCAGGCCAT ACGTCCGGTA	
36601				CAAGGCCAGC GTTCCGGTCG	
36651	GAACCGTAAA CTTGGCATTT	AAGGCCGCGT TTCCGGCGCA	TGCTGGCGTT ACGACCGCAA	TTTCCATAGG AAAGGTATCC	CTCCGCCCCC GAGGCGGGGG
36701	CTGACGAGCA GACTGCTCGT			GTCAGAGGTG CAGTCTCCAC	
36751	ACAGGACTAT TGTCCTGATA			CCTGGAAGCT GGACCTTCGA	
36801	CTCTCCTGTT GAGAGGACAA			ATACCTGTCC TATGGACAGG	
36851	CTTCGGGAAG GAAGCCCTTC			CACGCTGTAG GTGCGACATC	
36901	TCGGTGTAGG AGCCACATCC			TGTGTGCACG ACACACGTGC	

Figure 26 AM

37001		CGACTTATCG GCTGAATAGC			
37051		GGTATGTAGG CCATACATCC			
37101		TACACTAGAA ATGTGATCTT			
37151		CTTCGGAAAA GAAGCCTTTT			
37201		GTAGCGGTGG CATCGCCACC			
37251		GGATCTCAAG CCTAGAGTTC			
37301		GAACGAAAAC CTTGCTTTTG			
37351		TCTTCACCTA AGAAGTGGAT			
37401		GGTCTGACAG CCAGACTGTC			
37451		TGTCTATTTC ACAGATAAAG			
37501		TACGATACGG ATGCTATGCC			
37551		GAGACCCACG CTCTGGGTGC			
37601		GGAAGGGCCG CCTTCCCGGC			
37651	CCTCCATCCA GGAGGTAGGT	GTCTATTAAT CAGATAATTA	TGTTGCCGGG ACAACGGCCC	AAGCTAGAGT TTCGATCTCA	AAGTAGTTCG TTCATCAAGC
37701	CCAGTTAATA GGTCAATTAT	GTTTGCGCAA CAAACGCGTT	CGTTGTTGCC GCAACAACGG	ATTGCTACAG TAACGATGTC	GCATCGTGGT CGTAGCACCA
37751	GTCACGCTCG CAGTGCGAGC	TCGTTTGGTA AGCAAACCAT	TGGCTTCATT ACCGAAGTAA	CAGCTCCGGT GTCGAGGCCA	TCCCAACGAT AGGGTTGCTA
37801	CAAGGCGAGT GTTCCGCTCA	TACATGATCC ATGTACTAGG	CCCATGTTGT GGGTACAACA	GCAAAAAAGC CGTTTTTTCG	GGTTAGCTCC CCAATCGAGG
37851	TTCGGTCCTC AAGCCAGGAG	CGATCGTTGT GCTAGCAACA	CAGAAGTAAG GTCTTCATTC	TTGGCCGCAG AACCGGCGTC	TGTTATCACT ACAATAGTGA

Figure 26 AN

37951	GATGCTTTTC			CCAAGTCATT GGTTCAGTAA	
38001	TGTATGCGGC	GACCGAGTTG	CTCTTGCCCG	GCGTCAACAC	GGGATAATAC
	ACATACGCCG	CTGGCTCAAC	GAGAACGGGC	CGCAGTTGTG	CCCTATTATG
38051				CATCATTGGA	
	GCGCGGTGTA	TCGTCTTGAA	ATTTTCACGA	GTAGTAACCT	TTTGCAAGAA
38101				TGTTGAGATC	
	GCCCCGCTTT	TGAGAGTTCC	TAGAATGGCG	ACAACTCTAG	GTCAAGCTAC
38151				GCATCTTTTA	
	ATTGGGTGAG	CACGTGGGTT	GACTAGAAGT	CGTAGAAAAT	GAAAGTGGTC
38201	CGTTTCTGGG	TGAGCAAAAA	CAGGAAGGCA	AAATGCCGCA	AAAAAGGGAA
	GCAAAGACCC	ACTCGTTTTT	GTCCTTCCGT	TTTACGGCGT	TTTTTCCCTT
38251				TACTCTTCCT	
	ATTCCCGCTG	TGCCTTTACA	ACTTATGAGT	ATGAGAAGGA	AAAAGTTATA
38301	TATTGAAGCA	TTTATCAGGG	TTATTGTCTC	ATGAGCGGAT	ACATATTTGA
	ATAACTTCGT	AAATAGTCCC	AATAACAGAG	TACTCGCCTA	TGTATAAACT
38351	ATGTATTTAG	AAAAATAAAC	AAATAGGGGT	TCCGCGCACA	TTTCCCCGAA
	TACATAAATC	TTTTTATTTG	TTTATCCCCA	AGGCGCGTGT	AAAGGGGCTT
38401	AAGTGCCACC				
	TTCACGGTGG	ACTGCAGATT	CTTTGGTAAT	AATAGTACTG	TAATTGGATA
38451	AAAAATAGGC	GTATCACGAG	GCCCTTTCGT	CTTCAAGAAT	TGGATCCGAA
	TTTTTATCCG	CATAGTGCTC	CGGGAAAGCA	GAAGTTCTTA	ACCTAGGCTT
		PacI			
38501	TTCTTAATTT	CTTAATTAA	(SEO ID NO	:32)	
70707	1101144111	CTIMITIM	, x	· ,	

38501 TTCTTAATTT CTTAATTAA (SEQ ID NO:32) AAGAATTAAA GAATTAATT (SEQ ID NO:33)

Figure 26 AD

1	CATCATCAAT GTAGTAGTTA			GAAGCCAATA CTTCGGTTAT	
51	GGGGTGGAGT CCCCACCTCA			TGGGAACGGG ACCCTTGCCC	
101				GTGTGGCGGA CACACCGCCT	
151				GTGTGCGCCG CACACGCGGC	
201				GATGTTGTAG CTACAACATC	
251				GGGAAAACTG CCCTTTTGAC	
301				TAGCGCGTAA ATCGCGCATT	
351				AGACTCGCCC TCTGAGCGGG	
401				TTGGCGTTTT AACCGCAAAA	
451				CATATCATAA GTATAGTATT	
501				TGTTGACATT ACAACTGTAA	
551					AGCCCATATA TCGGGTATAT
601					TGGCTGACCG ACCGACTGGC
651	CCCAACGACC GGGTTGCTGG	CCCGCCCATT GGGCGGGTAA	GACGTCAATA CTGCAGTTAT	ATGACGTATG TACTGCATAC	TTCCCATAGT AAGGGTATCA
701	AACGCCAATA TTGCGGTTAT	GGGACTTTCC CCCTGAAAGG	ATTGACGTCA TAACTGCAGT	ATGGGTGGAG TACCCACCTC	TATTTACGGT ATAAATGCCA
751					AAGTACGCCC TTCATGCGGG
801	CCTATTGACG GGATAACTGC	TCAATGACGG AGTTACTGCC	TAAATGGCCC ATTTACCGGG	GCCTGGCATT CGGACCGTAA	ATGCCCAGTA TACGGGTCAT

Figure 27A

901			CGGTTTTGGC GCCAAAACCG	
951			ATTTCCAAGT TAAAGGTTCA	
1001			AAAATCAACG TTTTAGTTGC	
1051			CAAATGGGCG GTTTACCCGC	
1101			TTTAGTGAAC AAATCACTTG	
1151			TCCATAGAAG AGGTATCTTC	
1201			ATTGGAACGC TAACCTTGCG	
1251			GCAAGTGGTC CGTTCACCAG	
1301		-	ATGAGGAGGG TACTCCTCCC	
1351			CGCAGTGGGC GCGTCACCCG	
1401	=		TCACCTCCTC AGTGGAGGAG	
1451			GCCCAGGAGG CGGGTCCTCC	
1501			GAGGCCCATG CTCCGGGTAC	
1551	CCTGTCCCAC GGACAGGGTG		AGAAGGGCGG TCTTCCCGCC	
1601	CCCAGAAGAG GGGTCTTCTC		CTGGACCTGT GACCTGGACA	
1651	TACTTCCCCG ATGAAGGGGC		CTACACCCCC GATGTGGGGG	
1701	CCTGACCTTC GGACTGGAAG		TCAAGCTGGT AGTTCGACCA	
1751	TGGAGGAGGC ACCTCCTCCG		GAGAACAACT CTCTTGTTGA	

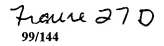
Figure 27B

1851	CTCCAAGCTG GAGGTTCGAC		GGAGCTGCAC CCTCGACGTG	
1901			CTGTGCCTTC GACACGGAAG	
1951			TCCTTGACCC AGGAACTGGG	
2001			GGAAATTGCA CCTTTAACGT	
2051			GGGTGGGGCA CCCACCCCGT	
2101			GCTGGGGATG CGACCCCTAC	
2151			TGTGGGCGTG ACACCCGCAC	
2201	•		GTAGTTTTGT CATCAAAACA	
2251			CGTTTGATGG GCAAACTACC	
2301			TGGGCCGGGG ACCCGGCCCC	
2351			CGTCCTGCCC GCAGGACGGG	
2401			CGCCGTTGGA GCGGCAACCT	
2451			GCCCGCGGA CGGGCGCCCT	
2501	CTTTGCTTTC GAAACGAAAG		TGCAGCTTCC ACGTCGAAGG	•
2551	CCCGCGATGA GGGCGCTACT		CACAATTGGA GTGTTAACCT	
2601	CGGGAACTTA GCCCTTGAAT		TTGGATCTGC AACCTAGACG	
2651	TTCTGCCCTG AAGACGGGAC			
2701	AAAAACCAGA TTTTTGGTCT		AGCAAGTGTC TCGTTCACAG	

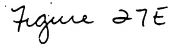
Figure 27C

2751	TATTTAGGGG	TTTTGCGCGC	GCGGTAGGCC	CGGGACCAGC	GGTCTCGGTC
2.02		AAAACGCGCG			
	MIMMICCCC	AAAACGCGCG	CGCCATCCGG	9000100	CCHOROCCHO
2801		CTGTGTATTT			
	CAACTCCCAG	GACACATAAA	AAAGGTCCTG	CACCATTTCC	ACTGAGACCT
2851	monnes es es	CATGGGCATA	* CCCCCTCTC	тесестесьс	GTAGCACCAC
2001					
	ACAAGTCTAT	GTACCCGTAT	TCGGGCAGAG	ACCCCACCTC	CATCGTGGTG
2901	TGCAGAGCTT	CATGCTGCGG	GGTGGTGTTG	TAGATGATCC	AGTCGTAGCA
=	ACCTCTCGAA	GTACGACGCC	CCACCACAAC	ATCTACTAGG	TCAGCATCGT
		••••••	•••••		

2951		GCGTGGTGCC			
	CCTCGCGACC	CGCACCACGG	ATTTTTACAG	AAAGTCATCG	TTCGACTAAC
3001	CCAGGGGCAG	GCCCTTGGTG	TAAGTGTTTA	CAAAGCGGTT	AAGCTGGGAT
5001		CGGGAACCAC			
	GGICCCCGIC	COGGAACCAC	ATTCACAAAT	G111COCC.2.	
3051	GGGTGCATAC	GTGGGGATAT	GAGATGCATC	TTGGACTGTA	TTTTTAGGTT
	CCCACGTATG	CACCCCTATA	CTCTACGTAG	AACCTGACAT	AAAAATCCAA
3101	CCCTATGTTC	CCAGCCATAT	CCCTCCGGGG	ATTCATGTTG	TGCAGAACCA
3101		GGTCGGTATA			
	CCGATACAAG	GGICGGIAIA	GGGAGGCCCC	IMGIACAAC	neorer root
3151	CCAGCACAGT	GTATCCGGTG	CACTTGGGAA	ATTTGTCATG	TAGCTTAGAA
	GGTCGTGTCA	CATAGGCCAC	GTGAACCCTT	TAAACAGTAC	ATCGAATCTT
3201	GGAAATGCGT	GGAAGAACTT	GGAGACGCCC	TTGTGACCTC	CAAGATTTTC
		CCTTCTTGAA			
	CCITIACGCA	cciiciida			
		TCCATAATGA	macca a macca	CCCACCCCC	CCCCCCTCCC
3251					
	GTACGTAAGC	AGGTATTACT	ACCGTTACCC	GGGTGCCCGC	CGCCGGACCC
3301	CGAAGATATT	TCTGGGATCA	CTAACGTCAT	AGTTGTGTTC	CAGGATGAGA
•••		AGACCCTAGT			
	GCIICINIAN	Moucecoulor			
		CCATTTTTAC	****	CCCACCCTCC	CACACTCCCC
3351					
	AGCAGTATCC	GGTAAAAATG	TTTCGCGCCC	GCCTCCCACG	GTCTGACGCC
3401	TATAATGGTT	CCATCCGGCC	CAGGGGCGTA	GTTACCCTCA	CAGATTTGCA
	ATATTACCAA	GGTAGGCCGG	GTCCCCGCAT	CAATGGGAGT	GTCTAAACGT
	717717 37700.27	••••••	••••		
2451	mmmaaaaaaa	mmmc » Cmmc »	CATCCCCCA	ייראייניירייאר	CTGCGGGGCG
3451	TTTCCCACGC	TITGAGTICA	GAIGGGGGA	1CAIGICIAC	C10CGGGGGG
	AAAGGGTGCG	AAACTCAAGT	CTACCCCCCT	AGTACAGATG	GACGCCCCGC
3501	ATGAAGAAAA	CGGTTTCCGG	GGTAGGGGAG	ATCAGCTGGG	AAGAAAGCAG
	ጥ ጀርጥጥር ጥ ጥጥ	GCCAAAGGCC	CCATCCCCTC	TAGTCGACCC	TTCTTTCGTC
	2770110111	••••			
2555	ommcooc: 65	************	MACCCC ACCC	CCTCCCCCC	TAAATCACAC
3551	GTTCCTGAGC	AGCTGCGACT	INCCOCNOCC	9919999999	17771 CACAC
	CAAGGACTCG	TCGACGCTGA	ATGGCGTCGG	CCACCCGGGC	ATTTAGTGTG
3601	CTATTACCGG	CTGCAACTGG	TAGTTAAGAG	AGCTGCAGCT	GCCGTCATCC
	GATAATGGCC	GACGTTGACC	ATCAATTCTC	TCGACGTCGA	CGGCAGTAGG
	J	2			
2655	000100100		COUNTY N CONTO	ጥርርርጥር እ ርጥር	GCATGTTTTC
3651	CTGAGCAGGG	GGGCCACTIC	GITANGCATG	1000107010	CCWIGITIE
	にない中では中でして	CCCGGTGAAG	CAATTCGTAC	AGGGACTGAG	CGTACAAAAG
	GACICGICCC				



3701				GCCCAGCGAT CGGGTCGCTA	
3751				GACCGTCCGC CTGGCAGGCG	
3801				CGGTCCCACA GCCAGGGTGT	
3851	·			TCCTCGTTTC AGGAGCAAAG	
3901				CTCGTCCAGA GAGCAGGTCT	
3951				TCAGCGTAGT AGTCGCATCA	
4001				GCCAGGGTGC CGGTCCCACG	
4051				TTCGCCCTGC AAGCGGGACG	
4101	CCATCGTAAA	CTGGTACCAC	AGTATCAGGT	GCCCCTCCGC CGGGGAGGCG	CCGCACCGGG
4151				CCGCACGAGG GGCGTGCTCC	
4201	TGAAAACTCC	CGCATCTCGA	ACCCGCGCTC	AAATACCGAT TTTATGGCTA	AGGCCCCTCA
4251	TCCGTAGGCG	CGGCGTCCGG	GGCGTCTGCC	AGAGCGTAAG	
4301	CACTCGAGAC	CGGCAAGCCC	CAGTTTTTGG	AGGTTTCCCC TCCAAAGGGG	GTACGAAAAA
		AATGGAGACC	AAAGGTACTC	GGCCACAGGT	GCGAGCCACT
		CAGGCACAGG	GGCATATGTC	TGAACTCTCC	GGACAGGAGC
		GCGCCAGGAG	GAGCATATCT	TTGAGCCTGG	TGAGACTCTG
		CAGGTCCGGT	CGTGCTTCCT	CCGATTCACC	CTCCCCATCG
		GTGATCCCCC	AGGTGAGCGA	GGTCCCACAC	TTCTGTGTAC
4601	TCGCCCTCTT AGCGGGAGAA		•	GGTTTGTAGG CCAAACATCC	



4701				CTGTTGGGGT GACAACCCCA
4751				GATTGTCAGT CTAACAGTCA
4801		GAGGAGGATT CTCCTCCTAA		GTGATGCCTT CACTACGGAA
4851		CGCATCCATC GCGTAGGTAG		
4901		CAAACGACCC GTTTGCTGGG		
4951		GTTTGGTTTT CAAACCAAAA		
5001		CACGTATTCG GTGCATAAGC		
5051		CGTCGGGCAC GCAGCCCGTG		
5101		TCAACGCTGG AGTTGCGACC		
5151		CCCCCCCCC		
5201		CGTCCGGGG GCAGGCCCCC		
5251		TCGAAGTAGT AGCTTCATCA		
5301		GCGGGCGGCA CGCCCGCCGT		GAGTGGGGGA CTCACCCCCT
5351	CCCCATGGCA GGGGTACCGT	TGGGGTGGGT ACCCCACCCA		
5401		AGGGGCTCTC TCCCCGAGAG		GGGTAGCATC CCCATCGTAG
5451	TTCCACCGCG AAGGTGGCGC			GTGCGAGGGA CACGCTCCCT
5501	GCGAGGAGGT CGCTCCTCCA	CGGGACCGAG GCCCTGGCTC		
5551				GTTGGACGCT CAACCTGCGA

Figure 27F

5651				AGCTCGGCGG TCGAGCCGCC	
5701				GATGATGTCA CTACTACAGT	
5751				GGACAAACTC CCTGTTTGAG	
5801				GCCTCCGAAC CGGAGGCTTG	
5851				GGCGCAGCAT CCGCGTCGTA	
5901				GGAGCGAGGT CCTCGCTCCA	
5951				TACTGGTATT ATGACCATAA	
6001				AAAGTCCGTG TTTCAGGCAC	
6051				CGTTGAAGAG GCAACTTCTC	
6101				AAGGGTCCCG TTCCCAGGGC	
6151				GATCTCGTCA CTAGAGCAGT	
6201				AGCGCGGGAT TCGCGCCCTA	
6251				AGCTCTTCAG TCGAGAAGTC	
6301					GAAGCGACGA CTTCGCTGCT
6351	ATGAGCTCCA TACTCGAGGT	CAGGTCACGG GTCCAGTGCC	GCCATTAGCA CGGTAATCGT	TTTGCAGGTG AAACGTCCAC	GTCGCGAAAG CAGCGCTTTC
6401	GTCCTAAACT CAGGATTTGA	GGCGACCTAT CCGCTGGATA	GGCCATTTTT CCGGTAAAAA	TCTGGGGTGA AGACCCCACT	TGCAGTAGAA ACGTCATCTT
6451	GGTAAGCGGG CCATTCGCCC	TCTTGTTCCC AGAACAAGGG	AGCGGTCCCA TCGCCAGGGT	TCCAAGGTTC AGGTTCCAAG	GCGGCTAGGT CGCCGATCCA
6501	CTCGCGCGGC GAGCGCGCCG	AGTCACTAGA TCAGTGATCT	GGCTCATCTC CCGAGTAGAG	CGCCGAACTT GCGGCTTGAA	CATGACCAGC GTACTGGTCG

Figure 276

6601		GTGACAAAGA CACTGTTTCT			
6651		GATCTCCCGC CTAGAGGGCG			
6701		AGTCCCTGCG TCAGGGACGC			
6751		CAGTACTGGC GTCATGACCG			
6801	GGTTGACCTG CCAACTGGAC	ACGACCGCGC TGCTGGCGCG			
6851		GGTTTGGCTG CCAAACCGAC			
6901		TGCTCGAGGG ACGAGCTCCC			
6951		AGTCCAGATG TCAGGTCTAC			
7001		GATGGGAGCT CTACCCTCGA			
7051	•	AGCTCCTGCA TCGAGGACGT			
7101		CAGGTGATAC GTCCACTATG			
7151	AGCTACCGAA	GCAAGAGGCC CGTTCTCCGG	CGTAGGGGCG	CCGCGCTGAT	GCCATGGCGC
7201	GCCGCCGCC	TGGGCCGCGG ACCCGGCGCC	CCCACAGGAA	CCTACTACGT	AGATTTTCGC
		GCTCGGGGGC	CTCCATCCCC	CCCGAGGCCT	GGGCGGCCCT
		CCCGTGCAGC	CGCGGCGCGC	GCCCGTCCTC	GACCACGACG
7351	GCGCGTAGGT CGCGCATCCA	TGCTGGCGAA ACGACCGCTT			
7401	GACCGCGGAG	TGCGTGAAGA ACGCACTTCT	GCTGCCCGGG	CCACTCGAAC	TTGGACTTTC
7451	AGAGTTCGAC TCTCAAGCTG	AGAATCAATT TCTTAGTTAA			

Figure 27H

7551		TCTTCCTCCT AGAAGGAGGA			
7601		GTCGTTGGAA CAGCAACCTT			
7651		CGTTCCAGAC GCAAGGTCTG			
7701		ATGACCACCT TACTGGTGGA			
7751		GTTTCGCAGG CAAAGCGTCC			
7801		CCACGAAGAA GGTGCTTCTT			
7851		CCCAAGGCCT GGGTTCCGGA			
7901		GAAAAACTGG CTTTTTGACC			
7951		GGATGAGCTC CCTACTCGAG			
8001	GGCTACAGGG CCGATGTCCC	GCCTCTTCTT CGGAGAAGAA	CTTCTTCAAT GAAGAAGTTA	CTCCTCTTCC GAGGAGAAGG	ATAAGGGCCT TATTCCCGGA
8051		TTCTTCTGGC AAGAAGACCG			
8101		CCGGGAGGCG GGCCCTCCGC			
8151		ATGGTCTCGG TACCAGAGCC			CGGGGGCGCA GCCCCCGCGT
8201	GTTGGAAGAC CAACCTTCTG	GCCGCCCGTC CGGCGGGCAG	ATGTCCCGGT TACAGGGCCA	TATGGGTTGG ATACCCAACC	CGGGGGGCTG GCCCCCGAC
8251	CCATGCGGCA GGTACGCCGT	GGGATACGGC CCCTATGCCG	GCTAACGATG CGATTGCTAC	CATCTCAACA GTAGAGTTGT	ATTGTTGTGT TAACAACACA
8301	AGGTACTCCG TCCATGAGGC	CCGCCGAGGG	ACCTGAGCGA TGGACTCGCT	GTCCGCATCG CAGGCGTAGC	ACCGGATCGG TGGCCTAGCC
8351	AAAACCTCTC TTTTGGAGAG	GAGAAAGGCG CTCTTTCCGC	TCTAACCAGT AGATTGGTCA	CACAGTCGCA GTGTCAGCGT	AGGTAGGCTG TCCATCCGAC
8401	AGCACCGTGG TCGTGGCACC	CGGGCGGCAG GCCCGCCGTC	CGGCGCCGCC	TCGGGGTTGT AGCCCCAACA	TTCTGGCGGA AAGACCGCCT

Figure 27I

8501				CCTGCTGAAT GGACGACTTA	
8551				CGGCGCAGGT GCCGCGTCCA	
8601				TTCTTCTCCT AAGAAGAGGA	
8651				CGGCGGAGTT GCCGCCTCAA	
8701				CCGAAGCCCC GGCTTCGGGG	
8751				GGCTAATATG CCGATTATAC	
8801				TGTCCACAAA ACAGGTGTTT	
8851				GCCATAACGG CGGTATTGCC	
8901	GGTCTGGTGA CCAGACCACT	CCCGGCTGCG GGGCCGACGC	AGAGCTCGGT TCTCGAGCCA	GTACCTGAGA CATGGACTCT	CGCGAGTAAG GCGCTCATTC
8951				TCCGCACCAG AGGCGTGGTC	
9001				TAGAGGGGCC ATCTCCCCGG	
9051	GGCCGGGCT CCGGCCCGA	CCGGGGGCGA GGCCCCCGCT	GATCTTCCAA CTAGAAGGTT	CATAAGGCGA GTATTCCGCT	TGATATCCGT ACTATAGGCA
9101				CGGCGGTGGT GCCGCCACCA	
9151	GGAAAGTCGC CCTTTCAGCG	GGACGCGGTT CCTGCGCCAA	CCAGATGTTG GGTCTACAAC	CGCAGCGGCA GCGTCGCCGT	AAAAGTGCTC TTTTCACGAG
9201	CATGGTCGGG GTACCAGCCC	ACGCTCTGGC TGCGAGACCG	CGGTCAGGCG GCCAGTCCGC	CGCGCAATCG GCGCGTTAGC	TTGACGCTCT AACTGCGAGA
9251	AGACCGTGCA TCTGGCACGT	AAAGGAGAGC TTTCCTCTCG	CTGTAAGCGG GACATTCGCC	GCACTCTTCC CGTGAGAAGG	GTGGTCTGGT CACCAGACCA
9301	GGATAAATTC CCTATTTAAG	GCAAGGGTAT CGTTCCCATA	CATGGCGGAC GTACCGCCTG	GACCGGGGTT CTGGCCCCAA	CGAGCCCCGT GCTCGGGGCA
9351	ATCCGGCCGT TAGGCCGGCA	CCGCCGTGAT GGCGGCACTA	CCATGCGGTT GGTACGCCAA	ACCGCCCGCG TGGCGGGGCGC	TGTCGAACCC ACAGCTTGGG

Figure 27J

9451		CTGCTGCGCT GACGACGCGA		GCGCGCAGCG CGCGCGTCGC
9501		GGCTGGAAAG CCGACCTTTC		
9551		ATTTTCCAAG TAAAAGGTTC		
9601		CGGACTGCGG GCCTGACGCC		
9651		GCAAATTCCT CGTTTAAGGA		
9701		GCATCCGGTG CGTAGGCCAC		
9751		AAGAGCAGCG TTCTCGTCGC		
9801		GGAGGGGCGA CCTCCCCGCT		
9851		CCCGCGCGCG		
9901		TGGCGCGGCT ACCGCGCCGA		
9951		AAGCGTGATA TTCGCACTAT		
10001		CCGCGAGGGA GGCGCTCCCT		
10051		GGCGCGAGCT CCGCGCTCGA		
10101	GCGCGAGGAG CGCGCTCCTC			AGTCCCGCGC TCAGGGCGCG
10151	GCGCACACGT CGCGTGTGCA	CCCCCCCCC		
10201	AACCAGGAGA TTGGTCCTCT			TGCGTACGCT ACGCATGCGA
10251	TGTGGCGCGC ACACCGCGCG	GAGGAGGTGG CTCCTCCACC		
	TAAGCGCGCT ATTCGCGCGA			

Figure 27K

10401				GCTGCTCGAT CGACGAGCTA	
10451				GCTTGAGCCT CGAAC,TCGGA	
10501				CTGGGCAAGT GACCCGTTCA	
10551				AGACAAGGAG TCTGTTCCTC	
10601				TGCTTACCTT ACGAATGGAA	
10651				AAGGCCGTGA TTCCGGCACT	
10701	GCGGCGCGAG CGCCGCGCTC	CTCAGCGACC GAGTCGCTGG	GCGAGCTGAT CGCTCGACTA	GCACAGCCTG CGTGTCGGAC	CAAAGGGCCC GTTTCCCGGG
10751	ACCGACCGTG	CCCGTCGCCG	CTATCTCTCC	CCGAGTCCTA GGCTCAGGAT	GAAACTGČGC
10801	CCGCGACTGG	ACGCGACCCG	GGGTTCGGCT	CGCGCCCTGG GCGCGGGACC	TCCGTCGACC
10851	CCGGCCTGGA	CCCGACCGCC	ACCGTGGGCG	GCGCGCTGGC CGCGCGACCG	TTGCAGCCGC
10901	CGCACCTCCT	TATACTGCTC	CTGCTACTCA	ACGAGCCAGA TGCTCGGTCT	CCTGCCGCTC
10951	ATGATTCGCC	ACTACAAAGA	CTAGTCTACT	TGCAAGACGC ACGTTCTGCG	TTGCCTGGGC
11001	CGCCACGCCC	GCCGCGACGT	CTCGGTCGGC	TCCGGCCTTA AGGCCGGAAT	TGAGGTGCCT
	GCTGACCGCG	GTCCAGTACC	TGGCGTAGTA	CAGCGACTGA	GCGCGCAATC CGCGCGTTAG
	GACTGCGCAA	GGCCGTCGTC	GGCGTCCGGT	TGGCCGAGAG	CGCAATTCTG GCGTTAAGAC
	CTTCGCCACC	AGGGCCGCGC	GCGTTTGGGG	TGCGTGCTCT	AGGTGCTGGC TCCACGACCG
	CTAGCATTTG	CGCGACCGGC	TTTTGTCCCG	GTAGGCCGGG	GACGAGGCCG CTGCTCCGGC
11251	GCCTGGTCTA CGGACCAGAT	CGACGCGCTG GCTGCGCGAC	CTTCAGCGCG GAAGTCGCGC	TGGCTCGTTA ACCGAGCAAT	CAACAGCGGC GTTGTCGCCG

Figure 27L

11351			CAACCTGGGC GTTGGACCCG	
11401			CCAACGTGCC GGTTGCACGG	
11451			CGGCTAATGG GCCGATTACC	
11501	· · · -		AGACTATTTT TCTGATAAAA	
11551			GCCAGGCTTT CGGTCCGAAA	
11601			GGCGACCGCG CCGCTGGCGC	
11.651			GCTGCTGCTA CGACGACGAT	
11701			CATACCTAGG GTATGGATCC	
11751			CATGTGGACG GTACACCTGC	
11801			GGGGCAGGAG CCCCGTCCTC	
11851			CCAACCGGCG GGTTGGCCGC	
11901			GAGCGCATTT CTCGCGTAAA	
11951			CGACGGGGTA GCTGCCCCAT	
12001	TGGCGCTGGA ACCGCGACCT		AACCGGGCAT TTGGCCCGTA	
12051	AACCGGCCGT TTGGCCGGCA		TACTTGCATC ATGAACGTAG	
12101	CGTGAACCCC GCACTTGGGG		CTTGAACCCG GAACTTGGGC	
12151	CGCCCCCTGG		AGGTGCCCGA TCCACGGGCT	
12201	GGATTCCTCT CCTAAGGAGA		GTGTTTTCCC CACAAAAGGG	

Figure 27 M

12301			CCGATCTAGG GGCTAGATCC	CGCTGCGGCC GCGACGCCGG
12351			AGCTTGATAG TCGAACTATC	
12401			GGGCGAGGAG CCCGCTCCTC	
12451			AAAACCTGCC TTTTGGACGG	
12501			AAGATGAGTA TTCTACTCAT	
12551			eeecececee cccececcce	
12601			TGTGGGAGGA ACACCCTCCT	
12651			GGGAGTGGCA CCCTCACCGT	
12701			TTAAAAAAAA TTTTTTTAA	
12751			GCACCGAGCG CGTGGCTCGC	
12801			ATGTATGAGG TACATACTCC	
12851			GCCAGTGGCG CGGTCACCGC	
12901			CGTTTGTGCC GCAAACACGG	
12951	CTGCGGCCTA GACGCCGGAT			AGTTGGCACC TCAACCGTGG
13001	CCTATTCGAC GGATAAGCTG			TCAACGGATG AGTTGCCTAC
13051	TGGCATCCCT ACCGTAGGGA			GACCACGGTC CTGGTGCCAG
13101	ATTCAAAACA TAAGTTTTGT			AGACCATCAA TCTGGTAGTT
	TCTTGACGAC AGAACTGCTG			

Figure 27N

13251		TGTCGCGCTT ACAGCGCGAA			
13301		GTGGAGTTCA CACCTCAAGT			
13351		CCTTATGAAC GGAATACTTG			
13401		ACGGGGTTCT TGCCCCAAGA			
13451		AGACTGGGGT TCTGACCCCA			
13501		AAACGAAGCC TTTGCTTCGG			
13551		ACTTCACCCA TGAAGTGGGT			
13601		CCCTTCCAGG GGGAAGGTCC			
13651		CATTCCCGCA GTAAGGGCGT			
13701		ACACCGAACA TGTGGCTTGT			
13751		GGCGCGGAAG CCGCGCCTTC			
13801		GGACATGAAC CCTGTACTTG			
13851		AGGAGAAGCG TCCTCTTCGC			
13901	CGCCCCCGCT GCGGGGGCGA	GCGCAACCCG CGCGTTGGGC	AGGTCGAGAA TCCAGCTCTT	GCCTCAGAAG CGGAGTCTTC	AAACCGGTGA TTTGGCCACT
13951	TCAAACCCCT AGTTTGGGGA	GACAGAGGAC CTGTCTCCTG	AGCAAGAAAC TCGTTCTTTG	GCAGTTACAA CGTCAATGTT	CCTAATAAGC GGATTATTCG
14001	AATGACAGCA TTACTGTCGT	CCTTCACCCA GGAAGTGGGT	GTACCGCAGC CATGGCGTCG	TGGTACCTTG ACCATGGAAC	CATACAACTA GTATGTTGAT
14051	CGGCGACCCT GCCGCTGGGA	CAGACCGGAA GTCTGGCCTT	TCCGCTCATG AGGCGAGTAC	GACCCTGCTT CTGGGACGAA	TGCACTCCTG ACGTGAGGAC
14101	ACGTAACCTG TGCATTGGAC	CGGCTCGGAG GCCGAGCCTC	CAGGTCTACT GTCCAGATGA	GGTCGTTGCC CCAGCAACGG	AGACATGATG TCTGTACTAC

Figure 270

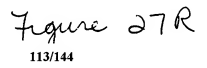
14201				TACAACGACC ATGTTGCTGG
14251	AGGCCGTCTA TCCGGCAGAT	CTCCCAACTC GAGGGTTGAG		
14301		TTCCCGAGAA AAGGGCTCTT		
14351		GTCAGTGAAA CAGTCACTTT		
14401		CAACAGCATC GTTGTCGTAG		CATTACTGAC GTAATGACTG
14451		GCACCTGCCC CGTGGACGGG		
14501		CTATCGAGCC GATAGCTCGG		
14551		CAATAACACA GTTATTGTGT	•	
14601		CCAAGAAGCG GGTTCTTCGC		
14651		GCGCCCTGGG CGCGGGACCC		
14701		TGACGCCATC ACTGCGGTAG		
14751		CGCCACCAGT GCGGTGGTCA	 	
14801		GCCCGGCGCT CGGGCCGCGA		
14851	TAGCACGTCG ATCGTGCAGC	CCACCGCCGC GGTGGCGGCG	 	
14901	GCGGCCCTGC CGCCGGGACG	TTAACCGCGC AATTGGCGCG		
14951	GGCCGCTCGA CCGGCGAGCT	AGGCTGGCCG TCCGACCGGC		
15001	GGCGACGAGC CCGCTGCTCG			
15051	GGTCGCAGGG CCAGCGTCCC			



15151	TGAATCTGAG	CATGACAACA	TACATAGGTC	CGCCGCCGC CGCCGCCGC	CGCGTTGCTT
15201	CGATACAGGT	TCGCGTTTTA	GTTTCTTCTC	ATGCTCCAGG TACGAGGTCC	AGTAGCGCGG
15251	CCTCTAGATA	CCGGGGGCT	TCTTCCTTCT	GCAGGATTAC CGTCCTAATG	TTCGGGGCTT
15301	TCGATTTCGC	CCAGTTTTTC	TTTTTCTTTC	ATGATGATGA TACTACTACT	ACTTGAACTG
15351	CTGCTCCACC	TTGACGACGT	GCGATGGCGC	CCCAGGCGAC GGGTCCGCTG	CCCATGTCAC
15401	CTTTCCAGCT	GCGCATTTTG	CACAAAACGC	ACCCGGCACC TGGGCCGTGG	TGGCATCAGA
15451	AATGCGGGCC	ACTCGCGAGG	TGGGCGTGGA	ACAAGCGCGT TGTTCGCGCA	CATACTACTC
15501	CACATGCCGC	TGCTCCTGGA	CGAACTCGTC	GCCAACGAGC CGGTTGCTCG	CGGAGCCCCT
15551	CAAACGGATG	CCTTTCGCCG	TATTCCTGTA	GCTGGCGTTG CGACCGCAAC	GGCGACCTGC
15601	TCCCGTTGGG	TTGTGGATCG	GATTTCGGGC	TAACACTGCA ATTGTGACGT	CGTCCACGAC
15651	GGGCGCGAAC	GTGGCAGGCT	TCTTTTCGCG	GGCCTAAAGC CCGGATTTCG	CGCTCAGACC
15701	ACTGAACCGT	GGGTGGCACG	TCGACTACCA	ACCCAAGCGC TGGGTTCGCG	GTCGCTGACC
15751	TTCTACAGAA	CCTTTTTTAC	TGGCACCTTG	CTGGGCTGGA GACCCGACCT	CGGGCTCCAG
		GTTAGTTCGT	CCACCGCGGC	CCTGACCCGC	ACGTCTGGCA
	CCTGCAAGTC	TATGGGTGAT	GGTCATCGTG	GTCATAACGG	
	TCCCGTACCT	CTGTGTTTGC	AGGGCCAAC	GGAGTCGCCA	
	CGCCACGTCC	GCCAGCGACG	CCGGCGCAGG	TTCTGGAGAT	
16001					CCGCGCCGTT

Figure 270

16051		CGCGCGCCCC			
16101		CGCCTACCCC GCGGATGGGG			
16151		ACTACCCGAC TGATGGGCTG			
16201		CCAGCCCGTG GGTCGGGCAC			
16251		GCAGGACCCT CGTCCTGGGA			
16301		AAGCCGGTCT TTCGGCCAGA			
16351		TTTCCCGGTG AAAGGGCCAC			
16401		CCGGCCACGG GGCCGGTGCC			
16451		CGCGCGTCGC GCGCGCAGCG			
16501		ACTGATCGCC TGACTAGCGG			
16551		TGCAGGCGCA ACGTCCGCGT			
16601		AATAAAAAGT TTATTTTCA			
16651		GAATGGAAGA CTTACCTTCT			CCCCGCGACA GGGGCGCTGT
16701	CGGCTCGCGC GCCGAGCGCG				ACCAGCAATA TGGTCGTTAT
16751	TGAGCGGTGG ACTCGCCACC	CGCCTTCAGC GCGGAAGTCG	TGGGGCTCGC ACCCCGAGCG	TGTGGAGCGG ACACCTCGCC	CATTAAAAAT GTAATTTTTA
16801	TTCGGTTCCA AAGCCAAGGT	CCGTTAAGAA GGCAATTCTT			
16851	AGGCCAGATG TCCGGTCTAC	CTGAGGGATA GACTCCCTAT	AGTTGAAAGA TCAACTTTCT	GCAAAATTTC CGTTTTAAAG	CAACAAAAGG GTTGTTTTCC
16901	TGGTAGATGG ACCATCTACC	CCTGGCCTCT GGACCGGAGA			
16951	CAGGCAGTGC GTCCGTCACG				GCCCTCCCGT CGGGAGGGCA



17051		GCGCCCCGAC CGCGGGGCTG		
17161		CGTACGAGGA GCATGCTCCT		
17151		CCCATGGCTA GGGTACCGAT		
17201	· - ·	GCCTCCCCC CGGAGGGGGG	= ' '	
17251		CCGTTGTTGT GGCAACAACA		
17301		GGTCCGCGAT CCAGGCGCTA		
17351		GAACAGCATC CTTGTCGTAG		
17401		TCTGATAGCT AGACTATCGA		
17451		CCAGAGGAGC GGTCTCCTCG		
17501		CTTCGATGAT GAAGCTACTA		
17551		TCGGAGTACC AGCCTCATGG		
17601		GTACTTCAGC CATGAAGTCG		
17651		ACGACGTGAC TGCTGCACTG		
17701	GTTCATCCCT CAAGTAGGGA	GTGGACCGTG CACCTGGCAC		
17751	TCACCCTAGC AGTGGGATCG	TGTGGGTGAT ACACCCACTA		
17801	TTTGACATCC AAACTGTAGG	GCGGCGTGCT CGCCGCACGA		
17851	TGGCACTGCC ACCGTGACGG	TACAACGCCC ATGTTGCGGG		
17901	AATGGGATGA TTACCCTACT	AGCTGCTACT TCGACGATGA		

Figure 275

17951				GCTGAGCAGC CGACTCGTCG	
18001		CAGGCGCCTT GTCCGCGGAA		AAATATTACA TTTATAATGT	
18051				AATATGCCGA	
		ACAGCTTCCA			ATTTTGTAAA
18101				TGGTACGAAA ACCATGCTTT	
18151				TACCCCAATG ATGGGGTTAC	
18201				ATGGAGGGCA TACCTCCCGT	
18251				CAAGTGGAAA GTTCACCTTT	
18301	CTCAACTACT GAGTTGATGA	CTCCGTCGGC	GTCCGTTACC	TGATAACTTG ACTATTGAAC	TGAGGATTTC
		010mc1101m	CMA CAMAMAC	AAACCCCAGA	ር እርጥር እጥ እጥጥ
18351		GTCACTTCTA			GTGAGTATAA
18401	ምር ጥጥ እ ር እ ሞርር	CCACTATTAA	CCAACCTAAC	TCACGAGAAC	TAATGGGCCA
18401		GGTGATAATT			ATTACCCGGT
18451	ACAATCTATG	CCCAACAGGC	CTAATTACAT	TGCTTTTAGG	GACAATTTTA
10151				ACGAAAATCC	
18501	TTGGTCTAAT	GTATTACAAC	AGCACGGGTA	ATATGGGTGT	TCTGGCGGGC
	AACCAGATTA	CATAATGTTG	TCGTGCCCAT	TATACCCACA	AGACCGCCCG
18551				TTGCAAGACA	
	GTTCGTAGCG	TCAACTTACG	ACAACATCTA	AACGTTCTGT	CTTTGTGTCT
18601	GCTTTCATAC	CAGCTTTTGC	TTGATTCCAT	TGGTGATAGA	ACCÁGGTACT
	CGAAAGTATG	GTCGAAAACG	AACTAAGGTA	ACCACTATCT	TGGTCCATGA
18651	TTTCTATGTG	GAATCAGGCT	GTTGACAGCT	ATGATCCAGA	TGTTAGAATT
	AAAGATACAC	CTTAGTCCGA	CAACTGTCGA	TACTAGGTCT	ACAATCTTAA
18701	ATTGAAAATC	ATGGAACTGA	AGATGAACTT	CCAAATTACT	GCTTTCCACT CGAAAGGTGA
18751	GGGAGGTGTG	ATTAATACAG	AGACTCTTAC	CAAGGTAAAA GTTCCATTTT	CCTAAAACAG
18801	GTCAGGAAAA Cagreerra	TGGATGGGAA	AAAGATGCTA TTTCTACGAT	CAGAATTTTC GTCTTAAAAG	AGATAAAAAT TCTATTTTTA
•					
18851	GAAATAAGAG CTTTATTCTC	TTGGAAATAA AACCTTTATT	TTTTGCCATG	GAAATCAATC CTTTAGTTAG	TAAATGCCAA ATTTACGGTT
	CITIMITCIC	- THE CALLES			

Figure 27T

18951		CAGTCCTTCC GTCAGGAAGG			
19001		TGAACAAGCG ACTTGTTCGC			
19051		GGAGCACGCT CCTCGTGCGA			
19101		CCACCGCAAT GGTGGCGTTA			
19151		GCTATGTGCC CGATACACGG			
19201		AACCTCCTTC TTGGAGGAAG			
19251		GGATGTTAAC CCTACAATTG			
19301		ACGGAGCCAG TGCCTCGGTC	•	GATAGCATTT CTATCGTAAA	
19351	GTGGAAGAAG	CCCATGGCCC GGGTACCGGG	TGTTGTGGCG	GAGGTGCGAA	CTCCGGTACG
19401	AATCTTTGCT	CACCAACGAC GTGGTTGCTG	GTCAGGAAAT	TGCTGATAGA	GAGGCGGCGG
19451	AACATGCTCT TTGTACGAGA	ACCCTATACC TGGGATATGG	CGCCAACGCT GCGGTTGCGA	ACCAACGTGC TGGTTGCACG	CCATATCCAT GGTATAGGTA
19501	GGGGAGGCG	AACTGGGCGG TTGACCCGCC	GAAAGGCGCC	GACCCGGAAG	TGCGCGGAAT
19551	AGACTAAGGA TCTGATTCCT	AACCCCATCA TTGGGGTAGT	CTGGGCTCGG GACCCGAGCC	GCTACGACCC CGATGCTGGG	TTATTACACC AATAATGTGG
19601	TACTCTGGCT ATGAGACCGA	CTATACCCTA GATATGGGAT	CCTAGATGGA GGATCTACCT	ACCTTTTACC TGGAAAATGG	TCAACCACAC AGTTGGTGTG
19651	CTTTAAGAAG GAAATTCTTC	GTGGCCATTA CACCGGTAAT	CCTTTGACTC GGAAACTGAG	TTCTGTCAGC AAGACAGTCG	TGGCCTGGCA ACCGGACCGT
19701	ATGACCGCCT TACTGGCGGA	GCTTACCCCC CGAATGGGGG	AACGAGTTTG TTGCTCAAAC	AAATTAAGCG TTTAATTCGC	CTCAGTTGAC GAGTCAACTG
19751	GGGGAGGGTT CCCCTCCCAA	ACAACGTTGC TGTTGCAACG	CCAGTGTAAC GGTCACATTG	ATGACCAAAG TACTGGTTTC	ACTGGTTCCT TGACCAAGGA
19801	GGTACAAATG CCATGTTTAC	CTAGCTAACT GATCGATTGA	ATAACATTGG TATTGTAACC	CTACCAGGGC GATGGTCCCG	TTCTATATCC AAGATATAGG

Figure 274

PCT/US01/28861 WO 02/022080

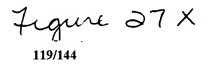
19851		CAAGGACCGC GTTCCTGGCG			
19901		AGGTGGTGGA TCCACCACCT			
19951		CACCAACACA GTGGTTGTGT			
20001		CGAAGGACAG GCTTCCTGTC			
20051		CCGCAGTTGA GGCGTCAACT			
20101		TGGCGCATCC ACCGCGTAGG			
20151		CCTGGGCCAA GGACCCGGTT			
20201		CTTTTGAGGT GAAAACTCCA			
20251		GAAGTCTTTG CTTCAGAAAC			
20301	CGCAGTAGCT	AACCGTGTAC TTGGCACATG	GACGCGTGCG	GGAAGAGCCG	GCCGTTGCGG
20351		GAAGCAAGCA CTTCGTTCGT			
20401	AGTGAGCAGG TCACTCGTCC	AACTGAAAGC TTGACTTTCG		GATCTTGGTT CTAGAACCAA	
20451		ACCTATGACA TGGATACTGT			
		GCGGTATCAG	TTATGCCGGC	CAGCGCTCTG	ACCCCCGCAT
		GGAAACGGAC	CTTGGGCGTG	AGTTTTTGTA	CGATGGAGAA
		CCGAAAAGAC	TGGTCGCTGA	GTTCGTCCAA	ATGGTCAAAC
		TGAGGACGCG	GCATCGCGGT	AACGAAGAAG	GGGGCTGGCG
20701	TGTATAACGC ACATATTGCG	TGGAAAAGTC ACCTTTTCAG			
20751	CGCCTGTGGA GCGGACACCT				GCCAACTGGC CGGTTGACCG

Figure 27 V.

20851		TGCTCAACAG ACGAGTTGTC			
20901		CTCTACAGCT GAGATGTCGA			
20951		GCAGATTAGG CGTCTAATCC			
21001		AATGTACTAG TTACATGATC			
21051		CTCGGGTGAT GAGCCCACTA			
21101	TTTAAAAATC	AAAGGGGTTC	TGCCGCGCAT	CGCTATGCGC	CACTGGCAGG
	AAATTTTTAG	TTTCCCCAAG	ACGGCGCGTA	GCGATACGCG	GTGACCGTCC
21151	GACACGTTGC	GATACTGGTG	TTTAGTGCTC	CACTTAAACT	CAGGCACAAC
	CTGTGCAACG	CTATGACCAC	AAATCACGAG	GTGAATTTGA	GTCCGTGTTG
21201	CATCCGCGGC	AGCTCGGTGA	AGTTTTCACT	CCACAGGCTG	CGCACCATCA
	GTAGGCGCCG	TCGAGCCACT	TCAAAAGTGA	GGTGTCCGAC	GCGTGGTAGT
21251	CCAACGCGTT	TAGCAGGTCG	GGCGCCGATA	TCTTGAAGTC	GCAGTTGGGG
	GGTTGCGCAA	ATCGTCCAGC	CCGCGGCTAT	AGAACTTCAG	CGTCAACCCC
21301	CCTCCGCCCT	GCGCGCGCGA	GTTGCGATAC	ACAGGGTTGC	AGCACTGGAA
	GGAGGCGGGA	CGCGCGCGCT	CAACGCTATG	TGTCCCAACG	TCGTGACCTT
21351	CACTATCAGC	GCCGGGTGGT	GCACGCTGGC	CAGCACGCTC	TTGTCGGAGA
	GTGATAGTCG	CGGCCCACCA	CGTGCGACCG	GTCGTGCGAG	AACAGCCTCT
21401	TCAGATCCGC	GTCCAGGTCC	TCCGCGTTGC	TCAGGGCGAA	CGGAGTCAAC
	AGTCTAGGCG	CAGGTCCAGG	AGGCGCAACG	AGTCCCGCTT	GCCTCAGTTG
21451	TTTGGTAGCT	GCCTTCCCAA	AAAGGGCGCG	TGCCCAGGCT	TTGAGTTGCA
	AAACCATCGA	CGGAAGGGTT	TTTCCCGCGC	ACGGGTCCGA	AACTCAACGT
21501	CTCGCACCGT	AGTGGCATCA	AAAGGTGACC	GTGCCCGGTC	TGGGCGTTAG
	GAGCGTGGCA	TCACCGTAGT	TTTCCACTGG	CACGGGCCAG	ACCCGCAATC
21551	GATACAGCGC	CTGCATAAAA	GCCTTGATCT	GCTTAAAAGC	CACCTGAGCC
	CTATGTCGCG	GACGTATTTT	CGGAACTAGA	CGAATTTTCG	GTGGACTCGG
21601	TTTGCGCCTT AAACGCGGAA	CAGAGAAGAA GTCTCTTCTT	CATGCCGCAA GTACGGCGTT	GACTTGCCGG	AAAACTGATT TTTTGACTAA
21651	GGCCGGACAG	GCCGCGTCGT	GCACGCAGCA	CCTTGCGTCG	GTGTTGGAGA
	CCGGCCTGTC	CGGCGCAGCA	CGTGCGTCGT	GGAACGCAGC	CACAACCTCT
21701	TCTGCACCAC	ATTTCGGCCC	CACCGGTTCT	TCACGATCTT	GGCCTTGCTA
	AGACGTGGTG	TAAAGCCGGG	GTGGCCAAGA	AGTGCTAGAA	CCGGAACGAT

Figure 27 W

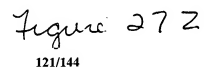
21801		TCCTTATTTA AGGAATAAAT		
21851		CTCAGCGCAG GAGTCGCGTC		
21901		TGTAGGTCAC ACATCCAGTG		
21951		ATCATCGTCA TAGTAGCAGT		
22001		GTGCTCCTCG CACGAGGAGC		
22051		GGTCAGGCAG CCAGTCCGTC	 	
22101		TTGTCCATCA AACAGGTAGT		
22151		GATCGGCACA CTAGCCGTGT	 	
22201		TGGGCTCTTC ACCCGAGAAG	 	
22251		TCTTCATTCA AGAAGTAAGT	 	
22301		TAGCACCGGT ATCGTGGCCA	 	
22351		TTTCTTCCTC AAAGAAGGAG		
22401		TTGGGAGAAG AACCCTCTTC		
22451	CCAAATCCGC GGTTTAGGCG	CGCCGAGGTC GCGGCTCCAG		
22501	AGCGCGTCTT TCGCGCAGAA	GTGATGAGTC CACTACTCAG		
	CATCCGCTTT GTAGGCGAAA			
22601	ACGACACGTC TGCTGTGCAG	CTCCATGGTT GAGGTACCAA		
22651	TCGGGGGTGG AGCCCCCACC			



22751			CCACCGATGC GGTGGCTACG	
22801			CTTGAGGAGG GAACTCCTCC	
22851			AGACGACGAG TCTGCTGCTC	
22901	••••	 	ACAACGCAGA TGTTGCGTCT	
22951			GGCGACTACC CCGCTGATGG	
23001			CCAGTGCGCC GGTCACGCGG	
23051			TCGCCATAGC AGCGGTATCG	
23101			CGCGTACCCC GCGCATGGGG	
23151			CCTCAACTTC GGAGTTGAAG	
23201			ACATCTTTTT TGTAGAAAAA	
23251			AGCCGAGCGG TCGGCTCGCC	
23301			TATCGCCTCG ATAGCGGAGC	
23351			ACGAGAAGCG TGCTCTTCGC	
23401	GCTCTGCAAC CGAGACGTTG			GAGTGTTGGT CTCACAACCA
23451	GGAACTCGAG CCTTGAGCTC		CGTACTAAAA GCATGATTTT	
23501	AGGTCACCCA TCCAGTGGGT		ACCTACCCCC TGGATGGGGG	
23551	AGCACAGTCA TCGTGTCAGT		CGTGCGCAGC GCACGCGTCG	
23601	GGATGCAAAT CCTACGTTTA			GCAGTTGGCG CGTCAACCGC

Figure 27 Y

23701				CTCGTTACCG GAGCAATGGC	
23751	CACGTACGTC	GCCAAGAAAC	GACTGGGCCT	GATGCAGCGC CTACGTCGCG	TTCGATCTCC
23801				ACGTACGCCA TGCATGCGGT	
23851	TAGAGGTTGC	ACCTCGAGAC	GTTGGACCAG	TCCTACCTTG AGGATGGAAC	CTTAAAACGT
23901	GCTTTTGGCG	GAACCCGTTT	TGCACGAAGT	TTCCACGCTC AAGGTGCGAG	TTCCCGCTCC
23951	GCGCGGCGCT	GATGCAGGCG	CTGACGCAAA	ACTTATTTCT TGAATAAAGA	TACGATGTGG
24001	ACCGTCTGCC	GGTACCCGCA	AACCGTCGTC	TGCTTGGAGG ACGAACCTCC	TCACGTTGGA
24051	GTTCCTCGAC	GTCTTTGACG	ATTTCGTTTT	CTTGAAGGAC GAACTTCCTG	GATACCTGCC
24101	GGAAGTTGCT	CGCGAGGCAC	CGGCGCGTGG	TGGCGGACAT ACCGCCTGTA	GTAAAAGGGG
24151	CTTGCGGACG	AATTTTGGGA	CGTTGTCCCA	CTGCCAGACT GACGGTCTGA	AGTGGTCAGT
24201	TTCGTACAAC	GTCTTGAAAT	CCTTGAAATA	CCTAGAGCGC GGATCTCGCG	AGTCCTTAGA
24251	ACGGGCGGTG	GACGACACGT	GAAGGATCGC	ACTTTGTGCC TGAAACACGG	GTAATTCATG
24301	GCGCTTACGG	GAGGCGGCGA	AACCCCGGTG	TGCTACCTTC ACGATGGAAG	ACGTCGATCG
		CGGATGGTGA	GACTGTATTA	CCTTCTGCAC	TCGCCACTGC
		CACAGTGACA	GCGACGTTGG	ATACGTGGGG	CGTGGCGAGG
		TAAGCGTCGA	CGAATTGCTT	TCAGTTTAAT	AGCCATGGAA
		CCAGGGAGCG	GACTGCTTTT	CAGGCGCCGA	GGCCCCAACT
24551	AACTCACTCC TTGAGTGAGG	GGGGCTGTGG	ACGTCGGCTT TGCAGCCGAA	ACCTTCGCAA TGGAAGCGTT	ATTTGTACCT TAAACATGGA



24601		ACGCCCACGA TGCGGGTGCT	 = '	-
24651		GAGCTTACCG CTCGAATGGC		
24701		AGCCATCAAC TCGGTAGTTG		
24751	-	TTTACTTGGA AAATGAACCT		
24801		CCGCAGCCCT GGCGTCGGGA		
24851	AGGATGGCAC TCCTACCGTG	CCAAAAAGAA GGTTTTTCTT		
24901		TGGGACAGTC ACCCTGTCAG		
24951		GGAAGACTGG CCTTCTGACC		
25001		CAGACGAAAC GTCTGCTTTG		
25051		AAATCGGCAA TTTAGCCGTT		
25101		GCCGGCACTG CGGCCGTGAC		
25151		CCAGGGCCGG GGTCCCGGCC		
25201		CAGCGCCAAG GTCGCGGTTC		
25251	CCATAGTTGC GGTATCAACG	TTGCTTGCAA AACGAACGTT		
25301	CGCTTTCTTC GCGAAAGAAG	TCTACCATCA AGATGGTAGT		
25351		CATCTCTACA GTAGAGATGT		
25401		CCACACAGAA GGTGTGTCTT		
25451	AAAGCCCAAG TTTCGGGTTC	AAATCCACAG TTTAGGTGTC		
25501	GTCTGGCGCC CAGACCGCGG			AAACAGGATT TTTGTCCTAA

Figure 27 AA

25551	TTTCCCACTC AAAGGGTGAG	TGTATGCTAT ACATACGATA	ATTTCAACAG TAAAGTTGTC	AGCAGGGGCC TCGTCCCCGG	AAGAACAAGA TTCTTGTTCT
25601				CCTCACCCGC GGAGTGGGCG	
25651				CGCTGGAAGA GCGACCTTCT	
25701				AAGGACTAGT TTCCTGATCA	
25751				TCTCCAGCGG AGAGGTCGCC	
25801	GCGGTCGTGG	ACAACAGTCG	CGGTAATACT	GCAAGGAAAT CGTTCCTTTA	AGGGTGCGGG
25851	ATGTACACCT	CAATGGTCGG	TGTTTACCCT	CTTGCGGCTG GAACGCCGAC	CTCGACGGGT
25901	TCTGATGAGT	TGGGCTTATT	TGATGTACTC	CGCGGGACCC GCGCCCTGGG	GTGTACTATA
25951	GGGCCCAGTT	GCCTTATGCG	CGGGTGGCTT	ACCGAATTCT TGGCTTAAGA	GGACCTTGTC
26001	CGCCGATAAT	GGTGGTGTGG	AGCATTATTG	CTTAATCCCC GAATTAGGGG	CATCAACCGG
26051	GCGACGGGAC	CACATGGTCC	TTTCAGGGCG	TCCCACCACT AGGGTGGTGA	CACCATGAAG
26101	GGTCTCTGCG	GGTCCGGCTT	CAAGTCTACT	CTAACTCAGG GATTGAGTCC	CCGCGTCGAA
26151	CGCCCGCCGA	AAGCAGTGTC	CCACGCCAGC	CCCGGGCAGG GGGCCCGTCC	CATATTGAGT
		TCTCCCGCTC	CATAAGTCGA	GTTGCTGCTC	AGCCACTCGA.
		AGAGGCAGGC	CTGCCCTGTA	AAGTCTAGCC	GCCGCGG CCG
		AGTGCGGAGC	AGTCCGTTAG	GATTGAGACG	TCTGGAGCAG
•		GCGAGACCTC	CGTAACCTTG	AGACGTTAAA	TAACTCCTCA
26401	TTGTGCCATC AACACGGTAG	GGTCTACTTT CCAGATGAAA	AACCCCTTCT	CGGGACCTCC	CGGCCACTAT
26451	CCGGATCAAT GGCCTAGTTA	TTATTCCTAA AATAAGGATT	CTTTGACGCG GAAACTGCGC	GTAAAGGACT CATTTCCTGA	CGGCGGACGG

Figure 27 AB

0 -, 0 0					101/050
26501	CTACGACTGA	ATGTTAAGTG	GAGAGGCAGA	GCAACTGCGC	CTGAAACACC
20301		TACAATTCAC			
	GATGUTGACT	TACAATTCAC	CICICCGICI	CG11GACGCG	GACTITO:00
26551	TGGTCCACTG	TCGCCGCCAC	AAGTGCTTTG	CCCGCGACTC	CGGTGAGTTT
		AGCGGCGGTG			
	ACCAGGTGAC	AGC GGC GG 1 G	TICACGAMAC	GGGCGC 1 GMG	GCCACTC/22.
26601	TGCTACTTTG	AATTGCCCGA	GGATCATATC	GAGGGCCCGG	CGCACGGCGT
		TTAACGGGCT			CCCTCCCCCA
	ACONTONANC	ITAACGGGCI	CCINGINING	6,6666666	000100000
			•		
26651	CCGGCTTACC	GCCCAGGGAG	AGCTTGCCCG	TAGCCTGATT	CGGGAGTTTA
	CCCCCAATCC	CGGGTCCCTC	TOGALACGOC	ATCGGACTAA	GCCCTCAAAT
	GGCCGWY1GG	6000166616	1 COMMCOOC	A1000	000010.
26701	CCCAGCGCCC	CCTGCTAGTT	GAGCGGGACA	GGGGACCCTG	TGTTCTCACT
	CCCTCCCCCC	GGACGATCAA	CTCGCCCTGT	CCCCTGGGAC	ACAAGAGTGA
	000100000	000			
26751		ACTGTCCTAA			
	CACTAAACGT	TGACAGGATT	GGGACCTAAT	GTAGTTCTAG	AAACAACGGT
	0.101121001			+	
					amacacamac
26801		GAGTATAATA			
	AGAGACACGA	CTCATATTAT	TTATGTCTTT	AATTTTATAT	GACCCCGAGG
			0.1.0000000000	******	CCNNNCCNNC
26851		CTGTAAACGC			
	ATAGCGGTAG	GACATTTGCG	GTGGCAGAAG	TGGGCGGGTT	CGTTTGGTTC
		CCTGGTACTT	mma a cameme	TCCCTCTCTCTC	אייייאראארא
26901					
	CGCTTGGAAT	GGACCATGAA	AATTGTAGAG	AGGGAGACAC	TAAATGTTGT
26051	C###C > > CCC	AGACGGAGTG	ACTOTACCAC	λαλλοοποπο	CGAGCTCAGC
26951	GITTCAACCC	AGACGGAGIG	AGICIACGAG	TOTALCICIC	COMCCACMCC
	CAAAGTTGGG	TCTGCCTCAC	TCAGATGCTC	TCTTGGAGAG	GCTCGAGTCG
27001	тастесатеа	GAAAAAACAC	CACCCTCCTT	ACCTGCCGGG	AACGTACGAG
27001	1701007107	CTTTTTTGTG	CMCCCACCAA	TOCACOCCC	TTCCATCCTC
	ATGAGGTAGT	CTTTTTTGTG	GIGGGAGGAA	IGGNCGGCCC	1100710010
27051	TGCGTCACCG	GCCGCTGCAC	CACACCTACC	GCCTGACCGT	AAACCAGACT
2.002		CGGCGACGTG			
	ACGCAG1 GGC	CGGCGACGIG	GIGIGGAIGG	COORCIOCEN	111001010
27101	TTTTCCGGAC	AGACCTCAAT	AACTCTGTTT	ACCAGAACAG	GAGGTGAGCT
	AAAAGGCCTG	TCTGGAGTTA	TTGAGACAAA	TGGTCTTGTC	CTCCACTCGA
	Mandoccio	10100;10111			
27151	TAGAAAACCC	TTAGGGTATT	AGGCCAAAGG	CGCAGCTACT	GTGGGGTTTA
	ATCTTTTGGG	AATCCCATAA	TCCGGTTTCC	GCGTCGATGA	CACCCCAAAT
				0m11mm0100	mmmcmcm t C t
27201	TGAACAATTC	AAGCAACTCT	ACGGGCTATT	CTAATTCAGG	TTTCTCTAGA
			- · · · ·		
	ACTTGTTAAG	TTCGTTGAGA	TGCCCGATAA	GATTAAGTCC	AAAGAGATCT
	ACTTGTTAAG	TTCGTTGAGA	TGCCCGATAA	GATTAAGTCC	AAAGAGATCT
	ACTTGTTAAG	TTCGTTGAGA	TGCCCGATAA	GATTAAGTCC	AAAGAGATCT
27251	ACTTGTTAAG ATCGGGGTTG	TTCGTTGAGA GGGTTATTCT	TGCCCGATAA CTGTCTTGTG	GATTAAGTCC ATTCTCTTTA	AAAGAGATCT TTCTTATACT
27251	ACTTGTTAAG ATCGGGGTTG	TTCGTTGAGA GGGTTATTCT	TGCCCGATAA CTGTCTTGTG	GATTAAGTCC ATTCTCTTTA	AAAGAGATCT
27251	ACTTGTTAAG ATCGGGGTTG	TTCGTTGAGA GGGTTATTCT	TGCCCGATAA CTGTCTTGTG	GATTAAGTCC ATTCTCTTTA	AAAGAGATCT TTCTTATACT
	ACTTGTTAAG ATCGGGGTTG TAGCCCCAAC	TTCGTTGAGA GGGTTATTCT CCCAATAAGA	TGCCCGATAA CTGTCTTGTG GACAGAACAC	GATTAAGTCC ATTCTCTTTA TAAGAGAAAT	AAAGAGATCT TTCTTATACT AAGAATATGA
	ACTTGTTAAG ATCGGGGTTG TAGCCCCAAC AACGCTTCTC	TTCGTTGAGA GGGTTATTCT CCCAATAAGA TGCCTAAGGC	TGCCCGATAA CTGTCTTGTG GACAGAACAC TCGCCGCCTG	GATTAAGTCC ATTCTCTTTA TAAGAGAAAT CTGTGTGCAC	AAAGAGATCT TTCTTATACT AAGAATATGA ATTTGCATTT
	ACTTGTTAAG ATCGGGGTTG TAGCCCCAAC AACGCTTCTC	TTCGTTGAGA GGGTTATTCT CCCAATAAGA TGCCTAAGGC	TGCCCGATAA CTGTCTTGTG GACAGAACAC TCGCCGCCTG	GATTAAGTCC ATTCTCTTTA TAAGAGAAAT CTGTGTGCAC	AAAGAGATCT TTCTTATACT AAGAATATGA
	ACTTGTTAAG ATCGGGGTTG TAGCCCCAAC AACGCTTCTC	TTCGTTGAGA GGGTTATTCT CCCAATAAGA TGCCTAAGGC	TGCCCGATAA CTGTCTTGTG GACAGAACAC TCGCCGCCTG	GATTAAGTCC ATTCTCTTTA TAAGAGAAAT CTGTGTGCAC	AAAGAGATCT TTCTTATACT AAGAATATGA ATTTGCATTT
27301	ACTTGTTAAG ATCGGGGTTG TAGCCCCAAC AACGCTTCTC TTGCGAAGAG	TTCGTTGAGA GGGTTATTCT CCCAATAAGA TGCCTAAGGC ACGGATTCCG	TGCCCGATAA CTGTCTTGTG GACAGAACAC TCGCCGCCTG AGCGGCGGAC	GATTAAGTCC ATTCTCTTTA TAAGAGAAAT CTGTGTGCAC GACACACGTG	AAAGAGATCT TTCTTATACT AAGAATATGA ATTTGCATTT TAAACGTAAA
27301	ACTTGTTAAG ATCGGGGTTG TAGCCCCAAC AACGCTTCTC TTGCGAAGAG ATTGTCAGCT	TTCGTTGAGA GGGTTATTCT CCCAATAAGA TGCCTAAGGC ACGGATTCCG TTTTAAACGC	TGCCCGATAA CTGTCTTGTG GACAGAACAC TCGCCGCCTG AGCGGCGGAC TGGGGTCGCC	GATTAAGTCC ATTCTCTTTA TAAGAGAAAT CTGTGTGCAC GACACACGTG ACCCAAGATG	AAAGAGATCT TTCTTATACT AAGAATATGA ATTTGCATTT TAAACGTAAA ATTAGGTACA
27301	ACTTGTTAAG ATCGGGGTTG TAGCCCCAAC AACGCTTCTC TTGCGAAGAG ATTGTCAGCT	TTCGTTGAGA GGGTTATTCT CCCAATAAGA TGCCTAAGGC ACGGATTCCG	TGCCCGATAA CTGTCTTGTG GACAGAACAC TCGCCGCCTG AGCGGCGGAC TGGGGTCGCC	GATTAAGTCC ATTCTCTTTA TAAGAGAAAT CTGTGTGCAC GACACACGTG ACCCAAGATG	AAAGAGATCT TTCTTATACT AAGAATATGA ATTTGCATTT TAAACGTAAA ATTAGGTACA
27301 27351	ACTTGTTAAG ATCGGGGTTG TAGCCCCAAC AACGCTTCTC TTGCGAAGAG ATTGTCAGCT TAACAGTCGA	TTCGTTGAGA GGGTTATTCT CCCAATAAGA TGCCTAAGGC ACGGATTCCG TTTTAAACGC AAAATTTGCG	TGCCCGATAA CTGTCTTGTG GACAGAACAC TCGCCGCCTG AGCGGCGGAC TGGGGTCGCC ACCCCAGCGG	GATTAAGTCC ATTCTCTTTA TAAGAGAAAT CTGTGTGCAC GACACACGTG ACCCAAGATG TGGGTTCTAC	AAAGAGATCT TTCTTATACT AAGAATATGA ATTTGCATTT TAAACGTAAA ATTAGGTACA TAATCCATGT
27301 27351	ACTTGTTAAG ATCGGGGTTG TAGCCCCAAC AACGCTTCTC TTGCGAAGAG ATTGTCAGCT TAACAGTCGA	TTCGTTGAGA GGGTTATTCT CCCAATAAGA TGCCTAAGGC ACGGATTCCG TTTTAAACGC AAAATTTGCG	TGCCCGATAA CTGTCTTGTG GACAGAACAC TCGCCGCCTG AGCGGCGGAC TGGGGTCGCC ACCCCAGCGG	GATTAAGTCC ATTCTCTTTA TAAGAGAAAT CTGTGTGCAC GACACACGTG ACCCAAGATG TGGGTTCTAC	AAAGAGATCT TTCTTATACT AAGAATATGA ATTTGCATTT TAAACGTAAA ATTAGGTACA TAATCCATGT
27301 27351	ACTTGTTAAG ATCGGGGTTG TAGCCCCAAC AACGCTTCTC TTGCGAAGAG ATTGTCAGCT TAACAGTCGA TAATCCTAGG	TTCGTTGAGA GGGTTATTCT CCCAATAAGA TGCCTAAGGC ACGGATTCCG TTTTAAACGC AAAATTTGCG TTTACTCACC	TGCCCGATAA CTGTCTTGTG GACAGAACAC TCGCCGCCTG AGCGGCGGAC TGGGGTCGCC ACCCCAGCGG CTTGCGTCAG	GATTAAGTCC ATTCTCTTTA TAAGAGAAAT CTGTGTGCAC GACACACGTG ACCCAAGATG TGGGTTCTAC CCCACGGTAC	AAAGAGATCT TTCTTATACT AAGAATATGA ATTTGCATTT TAAACGTAAA ATTAGGTACA TAATCCATGT CACCCAAAAG
27301 27351	ACTTGTTAAG ATCGGGGTTG TAGCCCCAAC AACGCTTCTC TTGCGAAGAG ATTGTCAGCT TAACAGTCGA TAATCCTAGG	TTCGTTGAGA GGGTTATTCT CCCAATAAGA TGCCTAAGGC ACGGATTCCG TTTTAAACGC AAAATTTGCG	TGCCCGATAA CTGTCTTGTG GACAGAACAC TCGCCGCCTG AGCGGCGGAC TGGGGTCGCC ACCCCAGCGG CTTGCGTCAG	GATTAAGTCC ATTCTCTTTA TAAGAGAAAT CTGTGTGCAC GACACACGTG ACCCAAGATG TGGGTTCTAC CCCACGGTAC	AAAGAGATCT TTCTTATACT AAGAATATGA ATTTGCATTT TAAACGTAAA ATTAGGTACA TAATCCATGT CACCCAAAAG

Figure 27AC

WO 02/022080 PCT/US01/28861 27451 GTGGATTTTA AGGAGCCAGC CTGTAATGTT ACATTCGCAG CTGAAGCTAA CACCTAAAAT TCCTCGGTCG GACATTACAA TGTAAGCGTC GACTTCGATT 27501 TGAGTGCACC ACTCTTATAA AATGCACCAC AGAACATGAA AAGCTGCTTA - ACTCACGTGG TGAGAATATT TTACGTGGTG TCTTGTACTT TTCGACGAAT 27551 TTCGCCACAA AAACAAAATT GGCAAGTATG CTGTTTATGC TATTTGGCAG AAGCGGTGTT TTTGTTTTAA CCGTTCATAC GACAAATACG ATAAACCGTC 27601 CCAGGTGACA CTACAGAGTA TAATGTTACA GTTTTCCAGG GTAAAAGTCA GGTCCACTGT GATGTCTCAT ATTACAATGT CAAAAGGTCC CATTTTCAGT 27651 TAAAACTTTT ATGTATACTT TTCCATTTTA TGAAATGTGC GACATTACCA ATTTTGAAAA TACATATGAA AAGGTAAAAT ACTTTACACG CTGTAATGGT 27701 TGTACATGAG CAAACAGTAT AAGTTGTGGC CCCCACAAAA TTGTGTGGAA ACATGTACTC GTTTGTCATA TTCAACACCG GGGGTGTTTT AACACACCTT 27751 AACACTGGCA CTTTCTGCTG CACTGCTATG CTAATTACAG TGCTCGCTTT TTGTGACCGT GAAAGACGAC GTGACGATAC GATTAATGTC ACGAGCGAAA 27801 GGTCTGTACC CTACTCTATA TTAAATACAA AAGCAGACGC AGCTTTATTG CCAGACATGG GATGAGATAT AATTTATGTT TTCGTCTGCG TCGAAATAAC 27851 AGGAAAAGAA AATGCCTTAA TTTACTAAGT TACAAAGCTA ATGTCACCAC TCCTTTTCTT TTACGGAATT AAATGATTCA ATGTTTCGAT TACAGTGGTG 27901 TAACTGCTTT ACTCGCTGCT TGCAAAACAA ATTCAAAAAG TTAGCATTAT ATTGACGAAA TGAGCGACGA ACGTTTTGTT TAAGTTTTTC AATCGTAATA 27951 AATTAGAATA GGATTTAAAC CCCCCGGTCA TTTCCTGCTC AATACCATTC TTAATCTTAT CCTAAATTTG GGGGGCCAGT AAAGGACGAG TTATGGTAAG 28001 CCCTGAACAA TTGACTCTAT GTGGGATATG CTCCAGCGCT ACAACCTTGA GGGACTTGTT AACTGAGATA CACCCTATAC GAGGTCGCGA TGTTGGAACT 28051 AGTCAGGCTT CCTGGATGTC AGCATCTGAC TTTGGCCAGC ACCTGTCCCG TCAGTCCGAA GGACCTACAG TCGTAGACTG AAACCGGTCG TGGACAGGGC 28101 CGGATTTGTT CCAGTCCAAC TACAGCGACC CACCCTAACA GAGATGACCA GCCTAAACAA GGTCAGGTTG ATGTCGCTGG GTGGGATTGT CTCTACTGGT 28151 ACACAACCAA CGCGGCCGCC GCTACCGGAC TTACATCTAC CACAAATACA TGTGTTGGTT GCGCCGGCGG CGATGGCCTG AATGTAGATG GTGTTTATGT 28201 CCCCAAGTTT CTGCCTTTGT CAATAACTGG GATAACTTGG GCATGTGGTG GGGGTTCAAA GACGGAAACA GTTATTGACC CTATTGAACC CGTACACCAC 28251 GTTCTCCATA GCGCTTATGT TTGTATGCCT TATTATTATG TGGCTCATCT CAAGAGGTAT CGCGAATACA AACATACGGA ATAATAATAC ACCGAGTAGA 28301 GCTGCCTAAA GCGCAAACGC GCCCGACCAC CCATCTATAG TCCCATCATT CGACGGATTT CGCGTTTGCG CGGGCTGGTG GGTAGATATC AGGGTAGTAA 28351 GTGCTACACC CAAACAATGA TGGAATCCAT AGATTGGACG GACTGAAACA

Figure 27AD

CACGATGTGG GTTTGTTACT ACCTTAGGTA TCTAACCTGC CTGACTTTGT

28451		CTGACCCTTG GACTGGGAAC			
28501		TCACATCGAA AGTGTAGCTT			
28551		GATTTGTCAC CTAAACAGTG			
28601		TTTATCCAGT AAATAGGTCA			
28651		CCATCCCCAG GGTAGGGGTC			
28701		AATTATGAAA TTAATACTTT			
28751		CGTTTTGTTC GCAAAACAAG			
28801		ACTCGTATAT TGAGCATATA			
28851		CGAAGCCTGG GCTTCGGACC			
28901		TCTTAGCCCT AGAATCGGGA			
28951	= '	GATGCCATGA CTACGGTACT			
29001		ACAAGTTGTT TGTTCAACAA			
29051	CGCCCACCTT GCGGGTGGAA	CTCCCACCCC GAGGGTGGGG	CACTGAAATC GTGACTTTAG	AGCTACTTTA TCGATGAAAT	ATCTAACAGG TAGATTGTCC
29101	AGGAGATGAC TCCTCTACTG				TATTACAGAG ATAATGTCTC
29151	CAGCGCCTGC GTCGCGGACG				AGCGCATGAA TCGCGTACTT
29201	TCAAGAGCTC AGTTCTCGAG				AGGGGTATCT TCCCCATAGA
	TTTGTCTCGT AAACAGAGCA				
29301	CACCGCCTTA GTGGCGGAAT				TGGTGGTCAT ACCACCAGTA

Figure 27 A E

29401				AGGATCTCTG TCCTAGAGAC	
29451				CCCTTTAACT GGGAAATTGA	
29501				TTAGCAAATT AATCGTTTAA	•
29551		- -	-	CAGCTCTGGT GTCGAGACCA	
29601				AAATGGAATG TTTACCTTAC	
29651	CCTGTTCCTG GGACAAGGAC			TCATGTTGTT AGTACAACAA	
29701				CCCGTGTATC GGGCACATAG	
29751				TACTCCTCCC ATGAGGAGGG	
29801				TACTCTCTTT ATGAGAGAAA	
29851				GCGCTCAAAA CGCGAGTTTT	
29901				CTCCCAAAAT GAGGGTTTTA	
29951				ACATAAACCT TGTATTTGGA	
30001	GCACCCCTCA CGTGGGGAGT			ACTGTGGCTG TGACACCGAC	
30051	TCTAATGGTC AGATTACCAG			GCAATCACAG CGTTAGTGTC	
30101	CCGTGCACGA GGCACGTGCT			CCCAAGGACC GGGTTCCTGG	
30151	TCAGAAGGAA AGTCTTCCTT			GGCCCCCTCA CCGGGGGAGT	
30201	TAGCAGTACC ATCGTCATGG				
30251	GTAGCTTGGG CATCGAACCC			TTTATACACA AAATATGTGT	

Lyure 27 AF

30351		GCAACTGGTC CGTTGACCAG		
30401		TACTGGAGCC ATGACCTCGG		
30451		CAGGAGGACT GTCCTCCTGA		
30501		AGTTATCCGT TCAATAGGCA		
30551		CCCTCTTTTT GGGAGAAAAA		
30601		GCCTTTACTT CGGAAATGAA		
30651		CTAAGCACTG GATTCGTGAC		
30701		TGCAGGAGAT ACGTCCTCTA		
30751		CCCTCAAAAC GGGAGTTTTG		
30801		ATGGTTCCTA TACCAAGGAT		
30851		TACAGTAGGA ATGTCATCCT		
30901		CTCCATCTCC GAGGTAGAGG		
30951		TTGGTCTTAA AACCAGAATT		
31001	TTTCAGTTTT AAAGTCAAAA			
31051	CAAAGTGCTC GTTTCACGAG	ATCTTATTAT TAGAATAATA		
31101	CAATTCCTTC GTTAAGGAAG	CTGGACCCAG GACCTGGGTC		
31151	CTGAAGGCAC GACTTCCGTG			
31201	GCTTATCCAA CGAATAGGTT	AATCTCACGG TTAGAGTGCC		

Figure 27 AG

31251			TGTAACACTA ACATTGTGAT	
31301			 CTCCAAGTGC GAGGTTCACG	
31351			 TACATTAATG ATGTAATTAC	
31401			 CCAAGAATAA GGTTCTTATT	
31451			 TTGCAGAAAA AACGTCTTTT	
31501			 ACATAGCTTA TGTATCGAAT	
31551	_		TATTCAACCT ATAAGTTGGA	
31601		0	 CCCCGGCTGG GGGGCCGACC	
31651			AGGTGTTATA TCCACAATAT	
31701			TATTAATAAA ATAATTATTT	
31751			TGCTGAGCCA ACGACTCGGT	
31801			AGGAGAAGTC TCCTCTTCAG	
31851			TAGGGCGGTG ATCCCGCCAC	
	AGCGCGCGAA TCGCGCGCTT			
31,951	CATGGCAGTG GTACCGTCAC			AGCATAAGGC TCGTATTCCG
32001	GCCTTGTCCT CGGAACAGGA			TAAATCAGCA ATTTAGTCGT
32051	CAGTAACTGC GTCATTGACG			CACAGTGCAA GTGTCACGTT
32101	GGCGCTGTAT CCGCGACATA			ACGTGGCCAT TGCACCGGTA
	CATACCACAA GTATGGTGTT			

Figure 27AH

32251		CTCTGATTAA GAGACTAATT		
32301		AACCTGCCCG TTGGACGGGC		
32351		AGTGGAGAGC TCACCTCTCG		
32401		TCAATGTTGG AGTTACAACC		
32451		AAGCTCCTCC TTCGAGGAGG	 	
32501		TCAGCGTAAA AGTCGCATTT		
32551		TGCATTGTCA ACGTAACAGT	 	
32601		GGTAGCGCGG CCATCGCGCC		
32651		GAGTGCGCCG CTCACGCGGC		
32701		GGAACGCCGG CCTTGCGGCC		
32751	-	GACAAACAGA CTGTTTGTCT		
32801		TAGTTGTAGT ATCAACATCA		
32851		GGGTTCTATG CCCAAGATAC	 	
32901	ACATCCACCA TGTAGGTGGT	CCGCAGAATA GGCGTCTTAT		
32951	CTGCGAGTCA GACGCTCAGT	CACACGGGAG GTGTGCCCTC		
33001	TTTTTTTTTT AAAAAAAAA	CCAAAAGATT GGTTTTCTAA		
33051		TCCCCTCCGG AGGGGAGGCC		
33101	AGATAATGGC TCTATTACCG	ATTTGTAAGA TAAACATTCT		

Figure 27 AI

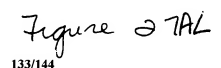
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33251	GCCACCTTCT	СААТАТАТСТ	CTAAGCAAAT	CCCGAATATT	AAGTCCGGCC
	CGGTGGAAGA	GTTATATAGA	GATTCGTTTA	GGGCTTATAA	TTCAGGCCGG
33301		TCTGCTCCAG AGACGAGGTC			
33351	AATCATGATT	GCAAAAATTC	AGGTTCCTCA	CAGACCTGTA	TAAGATTCAA
	TTAGTACTAA	CGTTTTTAAG	TCCAAGGAGT	GTCTGGACAT	ATTCTAAGTT
33401		TTAACAAAAA AATTGTTTTT			
33451		ATAATCGTGC TATTAGCACG			
33501		CCATGACAAA			
33301		GGTACTGTTT			
33551		CTAACCAGCG GATTGGTCGC			
33601		ATGCAAGGTG TACGTTCCAC			
33651		GCACATCGTA CGTGTAGCAT			
33701		ACCACAGAAA TGGTGTCTTT			
33751		CATAAACACA GTATTTGTGT			
33801		TCTTACAACA AGAATGTTGT			
33851		TGCCGGCGTG ACGGCCGCAC			
33901		GACAGCTCCT CTGTCGAGGA			
33951		ATCAGGTTGA TAGTCCAACT			
34001		GGGGAATACA CCCCTTATGT			
34051		GGTATAACAA CCATATTGTT			

Figure 27AJ

34151		CTTCCACAGC GAAGGTGTCG			
34201		CTATTAAAA GATAATTTTT		-	=
34251		TAAAAAAGGG ATTTTTTCCC			
34301		TAACGGTTAA ATTGCCAATT			
34351		GCCCAGAAAC CGGGTCTTTG			
34401	AGCAGTGAAG	CGTTTTCCCA GCAAAAGGGT	GCAATGCAGT	GAAGGGTAAA	ATTCTTTTGA
34451	TGTTAAGGGT	ACACATACAA TGTGTATGTT	CAATGAGGCG	GGATTTTGGA	TGCAGTGGGC
34501		ACGCCCCGCG TGCGGGGCGC			GGAGTAATAG
					PacI
34551		CAATCCAAAA GTTAGGTTTT			-
34601		TGCGACGCGA ACGCTGCGCT			
34651		GCGGCATCGG CGCCGTAGCC			
34701	CGTCCATCTA	GACGACCATC CTGCTGGTAG	TCCCTGTCGA	AGTTCCGGTC	GTTTTCCGGT
		TTTCCGGCGC	AACGACCGCA	AAAAGGTATC	CGAGGCGGGG
		TAGTGTTTTT	AGCTGCGAGT	TCAGTCTCCA	CCGCTTTGGG
		ATTTCTATGG	TCCGCAAAGG	GGGACCTTCG	AGGGAGCACG
34901	GCTCTCCTGT CGAGAGGACA	TCCGACCCTG AGGCTGGGAC			
34951	CCTTCGGGAA GGAAGCCCTT	GCGTGGCGCT CGCACCGCGA			
35001	TTCGGTGTAG AAGCCACATC	GTCGTTCGCT CAGCAAGCGA			

Figure 27 AK

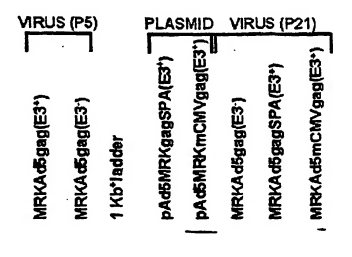
	AAGTCGGGCT	GGCGACGCGG	AATAGGCCAT	TGATAGCAGA	ACTCAGGTTG
35101	CCGCTAAGAC	ACGACTTATC	GCCACTGGCA	GCAGCCACTG	GTAACAGGAT
33101		TGCTGAATAG			
	GGCCATICIG	IGCIGAAIAG	CGGIGACCGI	CGICGGIGAC	CAITGICCIA
35151	TAGCAGAGCG	AGGTATGTAG	GCGGTGCTAC	AGAGTTCTTG	AAGTGGTGGC
	ATCGTCTCGC	TCCATACATC	CGCCACGATG	TCTCAAGAAC	TTCACCACCG
35201	CTAACTACGG	CTACACTAGA	AGGACAGTAT	TTGGTATCTG	CGCTCTGCTG
,0000		GATGTGATCT			
	GAT 1 GITTOCC	aniolanici	iccidicai.	DICCATAGAC	ocononconc
35251	AAGCCAGTTA	CCTTCGGAAA	AAGAGTTGGT	AGCTCTTGAT	CCGGCAAACA
		GGAAGCCTTT			•
35301	AACCACCGCT	GGTAGCGGTG	GTTTTTTTGT	TTGCAAGCAG	CAGATTACGC
		CCATCGCCAC			
			•••••		•======================================
35351	GCAGAAAAA	AGGATCTCAA	GAAGATCCTT	TGATCTTTTC	TACGGGGTCT
		TCCTAGAGTT			
	201011111	10011.01.01	C11C1CC.11		
35401	GACGCTCAGT	GGAACGAAAA	CTCACGTTAA	GGGATTTTGG	TCATGAGATT
	CTGCGAGTCA	CCTTGCTTTT	GAGTGCAATT	CCCTAAAACC	AGTACTCTAA
35451	ATCAAAAAGG	ATCTTCACCT	AGATCCTTTT	AAATCAATCT	AAAGTATATA
	TAGTTTTTCC	TAGAAGTGGA	TCTAGGAAAA	TTTAGTTAGA	TTTCATATAT
35501	TGAGTAAACT	TGGTCTGACA	GTTACCAATG	CTTAATCAGT	GAGGCACCTA
	ACTCATTTGA	ACCAGACTGT	CAATGGTTAC	GAATTAGTCA	CTCCGTGGAT
35551	TCTCAGCGAT	CTGTCTATTT	CGTTCATCCA	TAGTTGCCTG	ACTCCCCGTC
	AGAGTCGCTA	GACAGATAAA	GCAAGTAGGT	ATCAACGGAC	TGAGGGGCAG
35601	GTGTAGATAA	CTACGATACG	GGAGGGCTTA	CCATCTGGCC	CCAGTGCTGC
		GATGCTATGC			
35651	AATGATACCG	CGAGACCCAC	GCTCACCGGC	TCCAGATTTA	TCAGCAATAA
	TTACTATGGC	GCTCTGGGTG	CGAGTGGCCG	AGGTCTAAAT	AGTCGTTATT
					•
35701	ACCAGCCAGC	CGGAAGGGCC	GAGCGCAGAA	GTGGTCCTGC	AACTTTATCC
	TGGTCGGTCG	GCCTTCCCGG	CTCGCGTCTT	CACCAGGACG	TTGAAATAGG
	;				
35751	GCCTCCATCC	AGTCTATTAA	TTGTTGCCGG	GAAGCTAGAG	TAAGTAGTTC
	CGGAGGTAGG	TCAGATAATT	AACAACGGCC	CTTCGATCTC	ATTCATCAAG
35801	GCCAGTTAAT	AGTTTGCGCA	ACGTTGTTGC	CATTGCTACA	GGCATCGTGG
	CGGTCAATTA	TCAAACGCGT	TGCAACAACG	GTAACGATGT	CCGTAGCACC
35851	TGTCACGCTC	GTCGTTTGGT	ATGGCTTCAT	TCAGCTCCGG	TTCCCAACGA
		CAGCAAACCA			
35901	TCAAGGCGAG	TTACATGATC	CCCCATGTTG	TGCAAAAAAG	CGGTTAGCTC
	AGTTCCGCTC	AATGTACTAG	GGGGTACAAC	ACGTTTTTTC	GCCAATCGAG
35951	CTTCGGTCCT	CCGATCGTTG	TCAGAAGTAA	GTTGGCCGCA	GTGTTATCAC
					CACAATAGTG
	WE ICCCROOK				



36051		CTGTGACTGG			
	TCTACGAAAA	GACACTGACC	ACTCATGAGT	TGGTTCAGTA	AGACTCTTAT
36101	GTGTATGCGG	CGACCGAGTT	GCTCTTGCCC	GGCGTCAACA	CGGGATAATA
		GCTGGCTCAA			
			• • • • • • • • • • • • • • • • • • • •		
36151	CCGCGCCACA	TAGCAGAACT	TTAAAAGTGC	TCATCATTGG	AAAACGTTCT
		ATCGTCTTGA			
36201	TCGGGGCGAA	AACTCTCAAG	GATCTTACCG	CTGTTGAGAT	CCAGTTCGAT
••••	-	TTGAGAGTTC			
36251	GTAACCCACT	CGTGCACCCA	ACTGATCTTC	AGCATCTTTT	ACTTTCACCA
		GCACGTGGGT			
36301	GCGTTTCTGG	GTGAGCAAAA	ACAGGAAGGC	AAAATGCCGC	AAAAAAGGGA.
•••		CACTCGTTTT			
				· .	
36351	ATAAGGGCGA	CACGGAAATG	TTGAATACTC	ATACTCTTCC	TTTTTCAATA
		GTGCCTTTAC			
36401	TTATTGAAGC	ATTTATCAGG	GTTATTGTCT	CATGAGCGGA	TACATATTTG
	AATAACTTCG	TAAATAGTCC	CAATAACAGA	GTACTCGCCT	ATGTATAAAC
36451	AATGTATTTA	GAAAAATAAA	CAAATAGGGG	TTCCGCGCAC	ATTTCCCCGA
	TTACATAAAT	CTTTTTATTT	GTTTATCCCC	AAGGCGCGTG	TAAAGGGGCT
36501	AAAGTGCCAC	CTGACGTCTA	AGAAACCATT	ATTATCATGA	CATTAACCTA
	TTTCACGGTG	GACTGCAGAT	TCTTTGGTAA	TAATAGTACT	GTAATTGGAT
36551		CGTATCACGA			
	ATTTTTATCC	GCATAGTGCT	CCGGGAAAGC	AGAAGTTCTT	AACCTAGGCT
		PacI			

36601 ATTCTTAATT TCTTAATTAA (SEQ ID NO:34)
TAAGAATTAA AGAATTAATT (SEQ ID NO:35)

Figure 27 AM



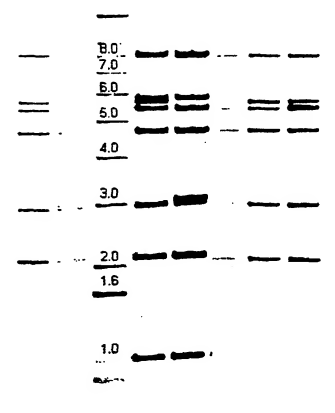


FIGURE 28

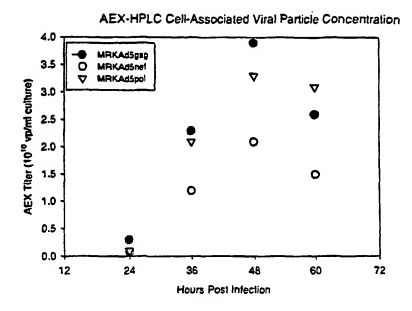


FIGURE 29A

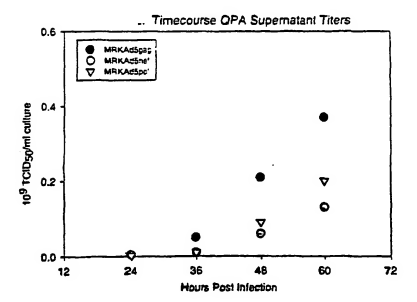


FIGURE 29B

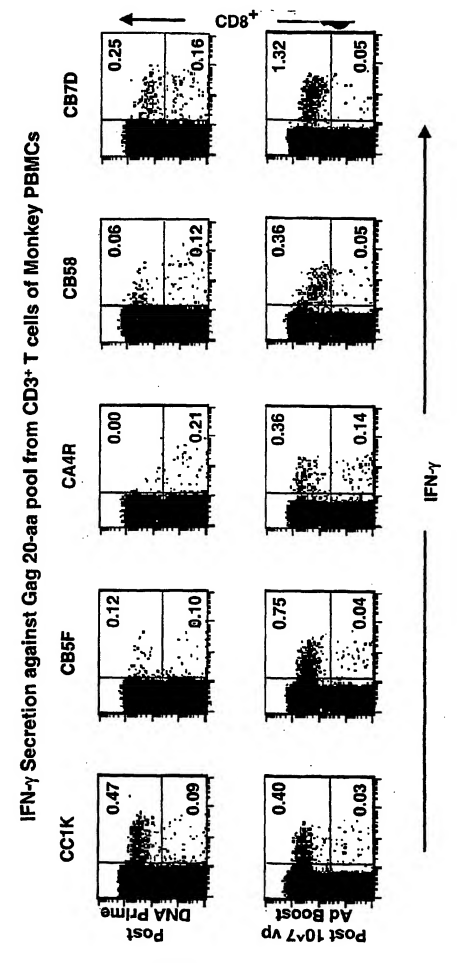
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gca Ala	gtc Val	ttc Phe	gtt Val 20	tcg Ser	ccc Pro	agc Ser	gag Glu	atc Ile 25	tcc Ser	att Ile	gtg Val	tgg Trp	gcc Ala 30	tcc Ser	agg Arg	96
gag Glu	ctg Leu	gag Glu 35	agg Arg	ttt Phe	gct Ala	gtg Val	aac Asn 40	cct Pro	ggc Gly	ctg Leu	ctg Leu	gag Glu 45	acc Thr	tct Ser	gag Glu	144
ggg ggg	tgc Cys 50	agg Arg	cag Gln	atc Ile	ctg Leu	ggc Gly 55	cag Gln	ctc Leu	cag Gln	ccc Pro	tcc Ser 60	ctg Leu	caa Gln	aca Thr	ggc	192
			ctg Leu													240
gtg Val	cac His	cag Gln	aag Lys	att Ile 85	gat Asp	gtg Val	aag Lys	gac Asp	acc Thr 90	aag Lys	gag Glu	gcc Ala	ctg Leu	gag Glu 95	aag Lys	288
att Ile	gag Glu	gag Glu	gag Glu 100	cag Gln	aac Asn	aag Lys	tcc Ser	aag Lys 105	aag Lys	aag Lys	gcc Ala	cag Gln	cag Gln 110	gct Ala	gct Ala	336
gct Ala	ggc Gly	aca Thr 115	Gly	aac Asn	tcc Ser	agc Ser	cag Gln 120	gtg Val	tcc Ser	cag Gln	aac Asn	tac Tyr 125	ccc Pro	att Ile	gtg Val	384
cag Gln	aac Asn 130	ctc Leu	cag Gln	Gly	cag Gln	atg Met 135	gtg Val	cac His	cag Gln	gcc Ala	atc Ile 140	tcc Ser	ccc Pro	cgg	acc	432
ctg Leu 145	Asn	gcc Ala	tgg Trp	gtg Val	aag Lys 150	gtg Val	gtg Val	gag Glu	gag Glu	aag Lys 155	gcc Ala	ttc Phe	tcc Ser	cct	gag Glu 160	480
gtg Val	atc Ile	ccc Pro	atg Met	ttc Phe 165	tct Ser	gcc Ala	ctg Leu	tct Ser	gag Glu 170	ggt Gly	gcc Ala	acc Thr	Pro	cag Gln 175	gac Asp	528
ctg Leu	aac Asn	acc	atg Met 180	ctg Leu	aac Asn	aca Thr	gtg Val	ggg Gly 185	ggc Gly	cat His	cag Gln	gct Ala	gcc Ala 190	Met	cag Gln	576
atg Met	ctg Leu	aag Lys 195	gag Glu	acc Thr	atc Ile	aat Asn	gag Glu 200	Glu	gct Ala	gct Ala	gag Glu	tgg Trp 205	Asp	agg Arg	ctg	624
cat His	Pro 210	Val	cac His	gct Ala	ggc	Pro 215	Ile	gcc	ccc Pro	ggc Gly	Cag Gln 220	Met	agg Arg	gag Glu	Pro	672
agg Arg 225	Gly	tct Ser	gac Asp	att Ile	gct Ala 230	ggc Gly	acc	acc	tcc Ser	acc Thr 235	Leu	cag Glr	gag Glu	cag Gln	att Ile 240	720
ggc Gly	tgg Trp	atg Met	acc Thr	aac Asn 245	Asn	Pro	Pro	atc	Pro 250	Val	ggg Gly	gaa Glu	atc Ile	tac Tyr 255	aag Lys	768

Figure 30'A"

agg Arg	tgg Trp	atc Ile	atc Ile 260	ctg Leu	ggc	ctg Leu	aac Asn	aag Lys 265	att Ile	gtg Val	agg Arg	atg Met	tac Tyr 270	tcc Ser	ccc Pro	816
acc	t <i>c</i> c Ser	atc Ile 275	ctg Leu	gac Asp	atc Ile	agg Ar g	cag Gln 280	ggc Gly	ccc Pro	aag Lys	gag Glu	ccc Pro 285	ttc Phe	agg Arg	gac Asp	864
													gcc Ala			912
													aat Asn			960
													gcc Ala			1008
gag Glu	gag Glu	atg Met	atg Met 340	aca Thr	gcc Ala	cys Cys	cag Gln	999 Gly 345	gtg Val	ej gaa	Gly	cct Pro	ggt Gly 350	cac His	aag Lys	1056
													tcc Ser			1104
													aca Thr			1152
	Phe												tgt Cys			1200
													cac His			1248
													atc Ile 430			1296
	His		Gly	Arg	Pro	Ğĺy		Phe		Gln	Ser		cct Pro			1344
aca Thr	gcc Ala 450	cct Pro	ccc Pro	gag Glu	gag Glu	tcc Ser 455	ttc Phe	agg Arg	ttť Phe	GJA 888	gag Glu 460	gag Glu	aag Lys	acc Thr	acc Thr	1392
ccc Pro 465	agc Ser	cag Gln	aag Lys	cag Gln	gag Glu 470	CCC Pro	att Ile	gac Asp	aag Lys	gag Glu 475	ctg Leu	tac Tyr	ccc Pro	ctg Leu	gcc Ala 480	1440
															D NO:36) D NO:37)	1482



Figure 31



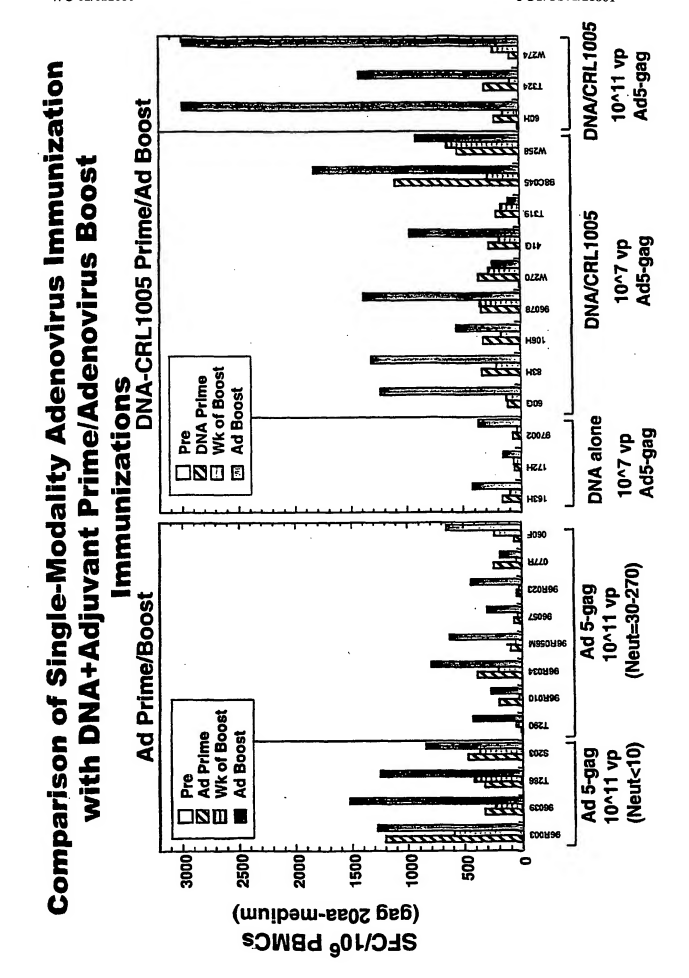


FIGURE 33A

ATGGGTGCTA	GGGCTTCTGT	GCTGTCTGGT	GGTGAGCTGG	ACAAGTGGGA	GAAGATCAGG
CTGAGGCCTG	GTGGCAAGAA	GAAGTACAAG	CTAAAGCACA	TTGTGTGGGC	CTCCAGGGAG
CTGGAGAGGT	TTGCTGTGAA	CCCTGGCCTG	CTGGAGACCT	CTGAGGGGTG	CAGGCAGATC
CTGGGCCAGC	TCCAGCCCTC	CCTGCAAACA	GGCTCTGAGG	AGCTGAGGTC	CCTGTACAAC
ACAGTGGCTA	CCCTGTACTG	TGTGCACCAG	AAGATTGATG	TGAAGGACAC	CAAGGAGGCC
CTGGAGAAGA	TTGAGGAGGA	GCAGAACAAG	TCCAAGAAGA	AGGCCCAGCA	GGCTGCTGCT
GGCACAGGCA	ACTCCAGCCA	GGTGTCCCAG	AACTACCCCA	TTGTGCAGAA	CCTCCAGGGC
CAGATGGTGC	ACCAGGCCAT	CTCCCCCGG	ACCCTGAATG	CCTGGGTGAA	GGTGGTGGAG
GAGAAGGCCT	TCTCCCCTGA	GGTGATCCCC	ATGTTCTCTG	CCCTGTCTGA	GGGTGCCACC
CCCCAGGACC	TGAACACCAT	GCTGAACACA	GTGGGGGGCC	ATCAGGCTGC	CATGCAGATG
CTGAAGGAGA	CCATCAATGA	GGAGGCTGCT	GAGTGGGACA	GGCTGCATCC	TGTGCACGCT
GGCCCCATTG	CCCCGGCCA	GATGAGGGAG	CCCAGGGGCT	CTGACATTGC	TGGCACCACC
TCCACCCTCC	AGGAGCAGAT	TGGCTGGATG	ACCAACAACC	CCCCCATCCC	TGTGGGGGAA
ATCTACAAGA	GGTGGATCAT	CCTGGGCCTG	AACAAGATTG	TGAGGATGTA	CTCCCCCACC
TCCATCCTGG	ACATCAGGCA	GGGCCCCAAG	GAGCCCTTCA	GGGACTATGT	GGACAGGTTC
TACAAGACCC	${\tt TGAGGGCTGA}$	GCAGGCCTCC	CAGGAGGTGA	AGAACTGGAT	GACAGAGACC
CTGCTGGTGC	AGAATGCCAA	CCCTGACTGC	AAGACCATCC	TGAAGGCCCT	GGGCCCTGCT
GCCACCCTGG	AGGAGATGAT	GACAGCCTGC	CAGGGGGTGG	GGGGCCCTGG	TCACAAGGCC
AGGGTGCTGG	CTGAGGCCAT	GTCCCAGGTG	ACCAACTCCG	CCACCATCAT	GATGCAGAGG
GGCAACTTCA	GGAACCAGAG	GAAGACAGTG	AAGTGCTTCA	ACTGTGGCAA	GGTGGGCCAC
ATTGCCAAGA	ACTGTAGGGC	CCCCAGGAAG	AAGGGCTGCT	GGAAGTGTGG	CAAGGAGGGC
CACCAGATGA	AGGACTGCAA	TGAGAGGCAG	GCCAACTTCC	TGGGCAAAAT	CTGGCCCTCC
CACAAGGGCA	GGCCTGGCAA	CTTCCTCCAG	TCCAGGCCTG	AGCCCACAGC	CCCTCCCGAG
	GGTTTGGGGA				
AAGGAGCTGT	ACCCCCTGGC	CTCCCTGAGG	TCCCTGTTTG	GCAACGACCC	CTCCTCCCAG
ATGGCTCCCA	TCTCCCCCAT	TGAGACTGTG	CCTGTGAAGC	TGAAGCCTGG	CATGGATGGC
	AGCAGTGGCC				
ACTGAGATGG	AGAAGGAGGG	CAAAATCTCC	AAGATTGGCC	CCGAGAACCC	CTACAACACC
-	CCATCAAGAA				
	AGAGGACCCA				
					CTTCTCTGTG
					CAACAATGAG
	TCAGGTACCA				
-					CCCTGACATT
	AGTACATGGC				
· · · · · · · · · · · · · · · · · · ·					CACCCTGAC
					CCCCGACAAG
					TGACATCCAG
- - ··					GGTGAGGCAG
					GACTGAGGAG
GCTGAGCTGG	AGCTGGCTGA	GAACAGGGAG	ATCCTGAAGG	AGCCTGTGCA	TGGGGTGTAC

FIGURE 33B

TATGACCCCT	CCAAGGACCT	GATTGCTGAG	ATCCAGAAGC	AGGGCCAGGG	CCAGTGGACC
TACCAAATCT	ACCAGGAGCC	CTTCAAGAAC	CTGAAGACTG	GCAAGTATGC	CAGGATGAGG
GGGGCCCACA	CCAATGATGT	GAAGCAGCTG	ACTGAGGCTG	TGCAGAAGAT	CACCACTGAG
TCCATTGTGA	TCTGGGGCAA	GACCCCCAAG	TTCAAGCTGC	CCATCCAGAA	GGAGACCTGG
GAGACCTGGT	GGACTGAGTA	CTGGCAGGCC	ACCTGGATCC	CTGAGTGGGA	GTTTGTGAAC
ACCCCCCCC	TGGTGAAGCT	GTGGTACCAG	CTGGAGAAGG	AGCCCATTGT	GGGGGCTGAG
ACCTTCTATG	TGGCTGGGGC	TGCCAACAGG	GAGACCAAGC	TGGGCAAGGC	TGGCTATGTG
ACCAACAGGG	GCAGGCAGAA	GGTGGTGACC	CTGACTGACA	CCACCAACCA	GAAGACTGCC
CTCCAGGCCA	TCTACCTGGC	CCTCCAGGAC	TCTGGCCTGG	AGGTGAACAT	TGTGACTGCC
TCCCAGTATG	CCCTGGGCAT	CATCCAGGCC	CAGCCTGATC	AGTCTGAGTC	TGAGCTGGTG
AACCAGATCA	TTGAGCAGCT	GATCAAGAAG	GAGAAGGTGT	ACCTGGCCTG	GGTGCCTGCC
CACAAGGGCA	TTGGGGGCAA	TGAGCAGGTG	GACAAGCTGG	TGTCTGCTGG	CATCAGGAAG
GTGCTGTTCC	TGGATGGCAT	TGACAAGGCC	CAGGATGAGC	ATGAGAAGTA	CCACTCCAAC
TGGAGGGCTA	TGGCCTCTGA	CTTCAACCTG	CCCCTGTGG	TGGCTAAGGA	GATTGTGGCC
TCCTGTGACA	AGTGCCAGCT	GAAGGGGGAG	GCCATGCATG	GGCAGGTGGA	CTGCTCCCCT
GGCATCTGGC	AGCTGGCCTG	CACCCACCTG	GAGGGCAAGG	TGATCCTGGT	GGCTGTGCAT
GTGGCCTCCG	GCTACATTGA	GGCTGAGGTG	ATCCCTGCTG	AGACAGGCCA	GGAGACTGCC
TACTTCCTGC	TGAAGCTGGC	TGGCAGGTGG	CCTGTGAAGA	CCATCCACAC	TGCCAATGGC
TCCAACTTCA	CTGGGGCCAC	AGTGAGGGCT	GCCTGCTGGT	GGGCTGGCAT	CAAGCAGGAG
TTTGGCATCC	CCTACAACCC	CCAGTCCCAG	GGGGTGGTGG	CCTCCATGAA	CAAGGAGCTG
AAGAAGATCA	TTGGGCAGGT	GAGGGACCAG	GCTGAGCACC	TGAAGACAGC	TGTGCAGATG
GCTGTGTTCA	TCCACAACTT	CAAGAGGAAG	GGGGGCATCG	GGGGCTACTC	CGCTGGGGAG
AGGATTGTGG	ACATCATTGC	CACAGACATC	CAGACCAAGG	AGCTCCAGAA	GCAGATCACC
AAGATCCAGA	ACTTCAGGGT	GTACTACAGG	GACTCCAGGA	ACCCCCTGTG	GAAGGGCCCT
-					TGACATCAAG
GTGGTGCCCA	GGAGGAAGGC	CAAGATCATC	AGGGACTATG	GCAAGCAGAT	GGCTGGGGAT
GACTGTGTGG	CCTCCAGGCA	GGATGAGGAC	TAA		
SEQ ID NO:	38				

FIGURE 34A

Met Gly Ala Arg Ala Ser Val Leu Ser Gly Gly Glu Leu Asp Lys Trp Glu Lys Ile Arg Leu Arg Pro Gly Gly Lys Lys Lys Tyr Lys Leu Lys His Ile Val Trp Ala Ser Arg Glu Leu Glu Arg Phe Ala Val Asn Pro Gly Leu Leu Glu Thr Ser Glu Gly Cys Arg Gln Ile Leu Gly Gln Leu Gln Pro Ser Leu Gln Thr Gly Ser Glu Glu Leu Arg Ser Leu Tyr Asn Thr Val Ala Thr Leu Tyr Cys Val His Gln Lys Ile Asp Val Lys Asp Thr Lys Glu Ala Leu Glu Lys Ile Glu Glu Glu Gln Asn Lys Ser Lys Lys Ala Gln Gln Ala Ala Ala Gly Thr Gly Asn Ser Ser Gln Val Ser Gln Asn Tyr Pro Ile Val Gln Asn Leu Gln Gly Gln Met Val His Gln Ala Ile Ser Pro Arg Thr Leu Asn Ala Trp Val Lys Val Val Glu Glu Lys Ala Phe Ser Pro Glu Val Ile Pro Met Phe Ser Ala Leu Ser Glu Gly Ala Thr Pro Gln Asp Leu Asn Thr Met Leu Asn Thr Val Gly Gly His Gln Ala Ala Met Gln Met Leu Lys Glu Thr Ile Asn Glu Glu Ala Ala Glu Trp Asp Arg Leu His Pro Val His Ala Gly Pro Ile Ala Pro Gly Gln Met Arg Glu Pro Arg Gly Ser Asp Ile Ala Gly Thr Thr Ser Thr Leu Gln Glu Gln Ile Gly Trp Met Thr Asn Asn Pro Pro Ile Pro Val Gly Glu Ile Tyr Lys Arg Trp Ile Ile Leu Gly Leu Asn Lys Ile Val Arg Met Tyr Ser Pro Thr Ser Ile Leu Asp Ile Arg Gln Gly Pro Lys Glu Pro Phe Arg Asp Tyr Val Asp Arg Phe Tyr Lys Thr Leu Arg Ala Glu Gln Ala Ser Gln Glu Val Lys Asn Trp Met Thr Glu Thr Leu Leu Val Gln Asn Ala Asn Pro Asp Cys Lys Thr Ile Leu Lys Ala Leu Gly Pro Ala Ala Thr Leu Glu Glu Met Met Thr Ala Cys Gln Gly Val Gly Pro Gly His Lys Ala Arg Val Leu Ala Glu Ala Met Ser Gln Val Thr Asn Ser Ala Thr Ile Met Met Gln Arg Gly Asn Phe Arg Asn Gln Arg Lys Thr Val Lys Cys Phe Asn Cys Gly Lys Val Gly His Ile Ala Lys Asn Cys Arg Ala Pro Arg Lys Lys Gly Cys Trp Lys Cys Gly Lys Glu Gly His Gln Met Lys Asp Cys Asn Glu Arg Gln Ala Asn Phe Leu Gly Lys Ile Trp Pro Ser His Lys Gly Arg Pro Gly Asn Phe Leu Gln Ser Arg Pro Glu Pro Thr Ala Pro Pro Glu Glu Ser Phe Arg Phe Gly Glu Glu Lys Thr Thr Pro Ser Gln Lys Gln Glu Pro Ile Asp Lys Glu Leu Tyr Pro Leu Ala Ser Leu Arg Ser Leu Phe Gly Asn Asp Pro Ser Ser Gln Met Ala Pro Ile Ser Pro Ile Glu Thr Val Pro Val Lys Leu Lys Pro Gly Met Asp Gly Pro Lys Val Lys Gln Trp Pro Leu Thr Glu Glu Lys Ile Lys Ala Leu Val Glu Ile Cys Thr Glu Met Glu Lys Glu Gly Lys Ile Ser Lys Ile Gly Pro Glu Asn Pro Tyr Asn Thr Pro Val Phe Ala Ile Lys Lys Lys Asp Ser Thr Lys Trp Arg Lys Leu Val Asp Phe Arg Glu Leu Asn Lys Arg Thr Gln Asp Phe Trp Glu Val Gln Leu Gly Ile Pro His Pro Ala Gly Leu Lys Lys Lys Ser Val Thr Val Leu Ala Val Gly Asp Ala Tyr Phe Ser Val Pro Leu Asp Glu Asp Phe Arg Lys Tyr Thr Ala Phe Thr Ile Pro Ser Ile Asn Asn Glu Thr Pro Gly Ile Arg Tyr Gln Tyr Asn Val Leu Pro Gln Gly Trp Lys Gly Ser Pro Ala Ile Phe Gln Ser Ser Met Thr Lys Ile Leu Glu Pro Phe Arg Lys Gln Asn Pro Asp Ile Val Ile Tyr Gln Tyr Met Ala Ala Leu Tyr Val Gly Ser Asp Leu Glu Ile Gly Gln His Arg Thr Lys Ile Glu Glu Leu Arg Gln His Leu Leu Arg Trp Gly Leu Thr Thr Pro Asp Lys Lys His Gln Lys Glu Pro Pro Phe Leu Trp Met Gly Tyr Glu Leu His Pro

FIGURE 34B

Asp Lys Trp Thr Val Gln Pro Ile Val Leu Pro Glu Lys Asp Ser Trp Thr Val Asn Asp Ile Gln Lys Leu Val Gly Lys Leu Asn Trp Ala Ser Gln Ile Tyr Pro Gly Ile Lys Val Arg Gln Leu Cys Lys Leu Leu Arg Gly Thr Lys Ala Leu Thr Glu Val Ile Pro Leu Thr Glu Glu Ala Glu Leu Glu Leu Ala Glu Asn Arg Glu Ile Leu Lys Glu Pro Val His Gly Val Tyr Tyr Asp Pro Ser Lys Asp Leu Ile Ala Glu Ile Gln Lys Gln Gly Gln Gly Gln Trp Thr Tyr Gln Ile Tyr Gln Glu Pro Phe Lys Asn Leu Lys Thr Gly Lys Tyr Ala Arg Met Arg Gly Ala His Thr Asn Asp Val Lys Gln Leu Thr Glu Ala Val Gln Lys Ile Thr Thr Glu Ser Ile Val Ile Trp Gly Lys Thr Pro Lys Phe Lys Leu Pro Ile Gln Lys Glu Thr Trp Glu Thr Trp Trp Thr Glu Tyr Trp Gln Ala Thr Trp Ile Pro Glu Trp Glu Phe Val Asn Thr Pro Pro Leu Val Lys Leu Trp Tyr Gln Leu Glu Lys Glu Pro Ile Val Gly Ala Glu Thr Phe Tyr Val Ala Gly Ala Ala Asn Arg Glu Thr Lys Leu Gly Lys Ala Gly Tyr Val Thr Asn Arg Gly Arg Gln Lys Val Val Thr Leu Thr Asp Thr Thr Asn Gln Lys Thr Ala Leu Gln Ala Ile Tyr Leu Ala Leu Gln Asp Ser Gly Leu Glu Val Asn Ile Val Thr Ala Ser Gln Tyr Ala Leu Gly Ile Ile Gln Ala Gln Pro Asp Gln Ser Glu Ser Glu Leu Val Asn Gln Ile Ile Glu Gln Leu Ile Lys Lys Glu Lys Val Tyr Leu Ala Trp Val Pro Ala His Lys Gly Ile Gly Gly Asn Glu Gln Val Asp Lys Leu Val Ser Ala Gly Ile Arg Lys Val Leu Phe Leu Asp Gly Ile Asp Lys Ala Gln Asp Glu His Glu Lys Tyr His Ser Asn Trp Arg Ala Met Ala Ser Asp Phe Asn Leu Pro Pro Val Val Ala Lys Glu Ile Val Ala Ser Cys Asp Lys Cys Gln Leu Lys Gly Glu Ala Met His Gly Gln Val Asp Cys Ser Pro Gly Ile Trp Gln Leu Ala Cys Thr His Leu Glu Gly Lys Val Ile Leu Val Ala Val His Val Ala Ser Gly Tyr Ile Glu Ala Glu Val Ile Pro Ala Glu Thr Gly Gln Glu Thr Ala Tyr Phe Leu Leu Lys Leu Ala Gly Arg Trp Pro Val Lys Thr Ile His Thr Ala Asn Gly Ser Asn Phe Thr Gly Ala Thr Val Arg Ala Ala Cys Trp Trp Ala Gly Ile Lys Gln Glu Phe Gly Ile Pro Tyr Asn Pro Gln Ser Gln Gly Val Val Ala Ser Met Asn Lys Glu Leu Lys Lys Ile Ile Gly Gln Val Arg Asp Gln Ala Glu His Leu Lys Thr Ala Val Gln Met Ala Val Phe Ile His Asn Phe Lys Arg Lys Gly Gly Jle Gly Gly Tyr Ser Ala Gly Glu Arg Ile Val Asp Ile Ile Ala Thr Asp Ile Gln Thr Lys Glu Leu Gln Lys Gln Ile Thr Lys Ile Gln Asn Phe Arg Val Tyr Tyr Arg Asp Ser Arg Asn Pro Leu Trp Lys Gly Pro Ala Lys Leu Leu Trp Lys Gly Glu Gly Ala Val Val Ile Gln Asp Asn Ser Asp Ile Lys Val Val Pro Arg Arg Lys Ala Lys Ile Ile Arg Asp Tyr Gly Lys Gln Met Ala Gly Asp Asp Cys Val Ala Ser Arg Gln Asp Glu Asp SEQ ID NO: 39

International application No.

PCT/US01/28861

		PC1/0501/	28801				
A. CLA	SSIFICATION OF SUBJECT MATTER						
IPC(7)	: C12N 15/86		1				
US CL	: 435/456		i				
	International Patent Classification (IPC) or to both r	ational classification and IPC					
B. FIEL	DS SEARCHED						
Minimum do	cumentation searched (classification system followed	by classification symbols)					
	24/205.1, 207.1, 227.1, 233.1; 435/69.1, 69.3, 173.						
	, , , , , , , , , , , , , , , , , , , ,	, ,					
Documentati	on searched other than minimum documentation to the	e extent that such documents are in	cluded in the fields searched				
	ta base consulted during the international search (nar	ne of data base and, where practice	able, search terms used)				
Please See C	ontinuation Sheet						
			ļ				
C. DOC	UMENTS CONSIDERED TO BE RELEVANT						
Category *	Citation of document, with indication, where ap	propriate, of the relevant passages	Relevant to claim No.				
X	WO 96/39178 (ERTL et al.) 12 December 1996 (12	<u> </u>					
	and claims 1 and 5.						
Y			4, 5, 13-17, 29-32, 34,				
_			35, 37				
x	US 6,019,978 A (ERTL et al.) 1 February 2000,(01	/02/2000), see columns 2, 7 and 8	1				
Y			4, 5, 13-17, 29-32, 34,				
			35, 37				
X,P	US 6,287,571 8 (ERTL et al.) 11 September 200	1 (11/09/2001), see columns 2, 7,	8 1, 9, 18				
	and claim 1.	., , , , , , , , , , , , , , , , , , ,	,,,,,,				
х	US 5,643,579A (HUNG et al.) 1 July 1997 (01/07/	1997), see examples 1, 2, 25 and 2	26. 1-3, 8, 9-11, 18				
	.,,	····,, ····· <u>··</u> ························					
Y			4,5,13-17, 29-32, 34,				
			35, 37				
Y	WANG et al. The use of an E1-deleted, replication	-defective adenovirus recombinant	1-3, 9-11, 13-18				
	expressing the rabies virus glycoprotein for early vi	accination of mice against rabies vi	irus.				
	Journal of Virology (March 1997) Vol. 71, No. 5,	р 3677-3683.					
Further	documents are listed in the continuation of Box C.	See patent family annex	ι.				
	pecial categories of cited documents:	• •	er the international filing date or				
			lict with the application but cited to				
"A" document	defining the general state of the art which is not considered to ticular relevance		heory underlying the invention				
oc or par	ICHIM ICICAMICS	"X" document of particular releva	ance; the claimed invention cannot be				
	plication or patent published on or after the international filing	considered novel or cannot b	e considered to involve an inventive				
date		step when the document is ta	uxen alone				
	which may throw doubts on priority claim(s) or which is cited		ance; the claimed invention cannot be				
to establi (as speci	sh the publication date of another citation or other special reason	considered to involve an invocation combined with one or more	entive step when the document is				
•		combination being obvious t					
O document	ent referring to an oral disclosure, use, exhibition or other means						
"P" document	"&" document member of the same patent family document published prior to the international filing date but later than the						
priority (late claimed	Date of malling of the land					
Date of the a	ctual completion of the international search	Date of mailing of the internation	nal search report				
06 February	2002 (06.02.2002)	19 AUG 200 2 00					
	ailing address of the ISA/US	Authorized officer	11 110				
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Box	PCT	Ulrike Winkler, Ph.D.	~~~ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\				
	hington, D.C. 20231	Telephone No. 703-308-0196	<i>[</i>]] [
· acomme M	o. (703)305-3230	101-300-0130					

Form PCT/ISA/210 (second sheet) (July 1998)

International application No.

PCT/US01/28861

INTERNATIONAL SEARCH REPORT

Category •	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No
Y	NATUK et al. Immunogenicity of recombinant human adenovirus -human immunodeficiency virus vaccines in chimpanzees. Aids Research and Human Retroviruses (1993) Vol. 9, No. 5, pp395-404, see material and methods.	1, 9, 29-32
Y	PREVEC et al. Immune response to HIV-1 gag antigens induced by recombinant adenovirus vectors in mice and rhesus macaque monkeys. Journal of Acquired Immune Deficincy Syndrome. (1991) Vol. 4, No. 6 pp. 568-76, see abstract.	1, 9, 29-32
Y	LORI et al. Rapid protection against human immunodeficiency virus type 1 (HIV-1) replication mediated by high efficiency non-retroviral delivery of genes interfering with HIV-1 tat and gag. Gene Therapy (1994) Vol. 1, No. 1, pp. 27-31, see abstract.	1, 9
Y	PFARR et al. Differential effects of polyadenylation regions on gene expression in mammalian cells. DNA (1986) Vol. 5, No. 2, pp.115-22, see abstract.	16
Y	NATUK et al. Adenovirus vectored vaccine. Developmental Biological Standards (1994) Vol. 82, pp. 71-77, see abstract.	1, 9

International application No.

PCT/US01/28861

Box	I Obse	ervations where certain claims were found unsearchable (Continuation of Item 1 of first sheet)						
This	interna	tional report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:						
1.		Claim Nos.: because they relate to subject matter not required to be searched by this Authority, namely:						
2.		Claim Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:						
3.	Claim Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).							
Box	II Ob	servations where unity of invention is lacking (Continuation of Item 2 of first sheet)						
		ional Searching Authority found multiple inventions in this international application, as follows: ontimuation Sheet						
1. 2. 3.		As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee. As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:						
4.	\boxtimes	No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.: 1-5, 8-11, 13-18, 29-32, 34, 35, 37						
Rem	iark on	Protest						

International application No.

PCT/US01/28861

BOX II. OBSERVATIONS WHERE UNITY OF INVENTION IS LACKING

This application contains the following inventions or groups of inventions which are not so linked as to form a single general inventive concept under PCT Rule 13.1. In order for all inventions to be examined, the appropriate additional examination fees must be paid.

Group	Claims	
1	1-5, 8-11,	The claims are directed to an adenoviral vector that is at least partially deleted of
-	13-18, 29,	ΔE1, the vector contains the cis-acting packaging sequence of the wild type
	30, 31, 32,	adenovirus genome, and a gene which encodes an HIV Gag protein (SEQ ID NO: 29)
	34, 35, 37	inserted in the parallel orientation of E1. In addition the vector contains a promoter
	1 .,,	and a polyadenylation signal.
2	6, 7, 36	The claims are directed to an adenoviral vector that is at least partially deleted of
-	1 0, ., 50	Δ E1 and Δ E3, the vector contains the cis-acting packaging sequence of the wild type
		adenovirus genome, and a gene which encodes an HIV Gag protein (SEQ ID NO:
		29).
3	12, 33	The claims are directed to an adenoviral vector that is at least partially deleted of
•	1 .2,00	ΔΕ1, the vector contains the cis-acting packaging sequence of the wild type
		adenovirus genome, and a gene which encodes an HIV protein inserted in the
		antiparallel orientation of E1.
4	19-23, 38-42	The claims are directed to a method of making and harvesting of a recombinant
•	1,7 23, 30 42	adenoviral particle that contains a gene encoding an HIV Gag protein.
5	24, 27, 28,	The claim is directed to a method of generating a cellular mediated immune response
•	43, 46, 47	to HIV Gag protein with the recombinant adenoviral particle.
6	25, 26, 44,	The claim is directed to a method of generating a cellular mediated immune response
v	45	to HIV Gag protein with the recombinant adenoviral particle in addition to
	~	administering a DNA plasmid vaccine.
7	48-51, 53,	The claims are directed to an adenoviral vector that is at least partially deleted of
′	54, 56	AE1, the vector contains the cis-acting packaging sequence of the wild type
	34, 30	adenovirus genome, and a gene which encodes an HIV Pol protein (SEQ ID NO: 1)
	•	inserted in the parallel orientation of E1.
8	48-51, 53,	The claims are directed to an adenoviral vector that is at least partially deleted of
• .	54, 56	
	34, 36	ΔΕ1, the vector contains the cis-acting packaging sequence of the wild type
		adenovirus genome, and a gene which encodes an HIV Pol protein (SEQ ID NO: 5)
9	48-51, 53,	inserted in the parallel orientation of E1. The claims are directed to an adenoviral vector that is at least partially deleted of
7	54, 56	
	34, 30	<u>AE1</u> , the vector contains the cis-acting packaging sequence of the wild type
		adenovirus genome, and a gene which encodes an HIV Pol protein (SEQ ID NO: 7)
10	- 62	inserted in the parallel orientation of E1.
10	52	The claim is directed to an adenoviral vector that is at least partially deleted of $\Delta E1$,
		the vector contains the cis-acting packaging sequence of the wild type adenovirus
		genome, and a gene which encodes an HIV Pol protein (SEQ ID NO: 1) inserted in
11		the antiparallel orientation of E1.
11	52	The claim is directed to an adenoviral vector that is at least partially deleted of $\Delta E1$,
		the vector contains the cis-acting packaging sequence of the wild type adenovirus
	-	genome, and a gene which encodes an HIV Pol protein (SEQ ID NO: 5) inserted in
10	 	the antiparallel orientation of E1.
12	52	The claim is directed to an adenoviral vector that is at least partially deleted of $\Delta E1$,
		the vector contains the cis-acting packaging sequence of the wild type adenovirus
		genome, and a gene which encodes an HIV Pol protein (SEQ ID NO: 7) inserted in
	+	the antiparallel orientation of E1.
13	55	The claim is directed to an adenoviral vector that is at least partially deleted of ΔΕ1

International application No.

PCT/US01/28861

		and $\Delta E3$, the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Pol protein (SEQ ID NO: 1) inserted in E1.
14	55	The claim is directed to an adenoviral vector that is at least partially deleted of $\Delta E1$ and $\Delta E3$, the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Pol protein (SEQ ID NO: 5) inserted in E1.
15	55	The claim is directed to an adenoviral vector that is at least partially deleted of $\Delta E1$ and $\Delta E3$, the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Pol protein (SEQ ID NO: 7) inserted in E1.
16	57-61	The claims are directed to a method of making and harvesting of a recombinant adenoviral particle that contains a gene encoding an HIV Pol protein.
17	62, 65, 66	The claim is directed to a method of generating a cellular mediated immune response to HIV Pol protein with the recombinant adenoviral particle.
18	63, 64	The claim is directed to a method of generating a cellular mediated immune response to HIV Pol protein with the recombinant adenoviral particle in addition to administering a DNA plasmid vaccine.
19	67-70, 72, 73, 75	The claims are directed to an adenoviral vector that is at least partially deleted of $\Delta E1$, the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Nef protein (SEQ ID NO: 9) inserted in the parallel orientation of E1.
20	67-70, 72, 73, 75	The claims are directed to an adenoviral vector that is at least partially deleted of $\Delta E1$, the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Nef protein (SEQ ID NO: 11) inserted in the parallel orientation of E1.
21	67-70, 72, 73, 75	The claims are directed to an adenoviral vector that is at least partially deleted of <u>AEI</u> , the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Nef protein (SEQ ID NO: 13) inserted in the parallel orientation of E1.
22	67-70, 72, 73, 75	The claims are directed to an adenoviral vector that is at least partially deleted of <u>\Delta E1</u> , the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an <u>HIV Nef protein (SEQ ID NO: 15)</u> inserted in the parallel orientation of E1.
23	71	The claim is directed to an adenoviral vector that is at least partially deleted of $\Delta E1$, the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Nef protein (SEQ ID NO: 9) inserted in the antiparallel orientation of E1.
24	71	The claim is directed to an adenoviral vector that is at least partially deleted of $\Delta E1$, the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Nef protein (SEQ ID NO: 11) inserted in the antiparallel orientation of E1.
25	71	The claim is directed to an adenoviral vector that is at least partially deleted of $\Delta E1$, the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Nef protein (SEQ ID NO: 13) inserted in the antiparallel orientation of E1.
26	71	The claim is directed to an adenoviral vector that is at least partially deleted of $\Delta E1$, the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Nef protein (SEQ ID NO: 15) inserted in the antiparallel orientation of E1.
27	74	The claim is directed to an adenoviral vector that is at least partially deleted of $\Delta E1$ and $\Delta E3$, the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Nef protein (SEQ ID NO: 9) inserted in E1.
28	74	The claim is directed to an adenoviral vector that is at least partially deleted of $\Delta E1$ and $\Delta E3$, the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Nef protein (SEQ ID NO: 11) inserted in E1.
29	74	The claim is directed to an adenoviral vector that is at least partially deleted of $\Delta E1$ and $\Delta E3$, the vector contains the cis-acting packaging sequence of the wild type

International application No.

PCT/US01/28861

		adenovirus genome, and a gene which encodes an HIV Nef protein (SEQ ID NO: 13) inserted in E1.
30	74	The claim is directed to an adenoviral vector that is at least partially deleted of $\Delta E1$ and $\Delta E3$, the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Nef protein (SEQ ID NO: 15) inserted in E1.
31	76-80	The claims are directed to a method of making and harvesting of a recombinant adenoviral particle that contains a gene encoding an HIV Nef protein.
32	81, 84, 85	The claims are directed to a method of generating a cellular mediated immune response to HIV Nef with the recombinant adenoviral particle.
33	82, 83	The claims are directed to a method of generating a cellular mediated immune response to HIV Nef with the recombinant adenoviral particle in addition to administering a DNA plasmid vaccine.
34	86a	The claim is drawn to a multivalent vaccine wherein gag, pol and nef are expressed from three individual vectors.
35	86b, 88, 89	The claims are drawn to a multivalent vaccine wherein gag, pol and nef are expressed from one individual vectors.
36	86c, 88	The claims are drawn to a multivalent vaccine wherein gag, pol and nef are expressed from two individual vectors, one expressing nef-pol fusion and one expressing gag.
37	86d, 87, 88	The claims are drawn to a multivalent vaccine wherein gag, pol and nef are expressed from two individual vectors, one expressing gag-pol fusion and one expressing nef.
38	86e, 88	The claims are drawn to a multivalent vaccine wherein gag, pol and nef are expressed from two individual vectors, one expressing nef-gag fusion and one expressing pol.
39	86f, 88	The claims are drawn to a multivalent vaccine wherein gag, pol and nef are expressed from a single vectors as a fusion protein.
40	86g, 88	The claims are drawn to a multivalent vaccine wherein gag and pol are expressed from two individual vectors.
41	86h, 88, 89	The claims are drawn to a multivalent vaccine wherein gag and pol are expressed individually from one vector.
42	86i, 88	The claims are drawn to a multivalent vaccine wherein pol and nef are expressed from two individual vectors.
43	86j, 88, 89	The claims are drawn to a multivalent vaccine wherein pol and nef are expressed from individually from one vector.
44	86k, 88	The claims are drawn to a multivalent vaccine wherein nef and gag are expressed individually from one vector.
45	861, 88, 89	The claims are drawn to a multivalent vaccine wherein nef and gag are expressed individually from one vector.
46	86m, 88	The claims are drawn to a multivalent vaccine wherein gag and pol are expressed as a fusion protein from one vector.
47	86n, 88	The claims are drawn to a multivalent vaccine wherein pol and nef are expressed as a fusion protein from one vector.
48	860, 88	The claims are drawn to a multivalent vaccine wherein nef and gag are expressed as a

The inventions listed as Groups 1-48 do not relate to a single general inventive concept under PCT Rule 13.1 because, under PCT Rule 13.2, they lack the same or corresponding special technical features for the following reasons:

The technical feature linking groups 1-33 appears to be a recombinant adenoviral vector wherein the adenoviral vector is at least partially deleted in E1 but the vector may contain more deletions, the vector contains wild type sequences including packaging signals and a gene encoding a heterologous HIV protein or fragments thereof. Ertl et al. (WO 96/39178) disclose a recombinant adenoviral vector that is deleted in E1 and partially deleted in E3, the remainder of the adenoviral vector contains wild type sequences. The vector additionally contains an insertion of a heterologous protein which includes HIV proteins (see abstract and claims 1 and 5). Therefore, the technical feature linking the inventions of groups 1-45 does not constitute a special technical feature as defined by PCT Rule 13.2, as it does not define a contribution over the prior art.

The special technical feature of the following groups 1-3, 7-15, 19-30 and 34-48 is considered to be the combination of sequences that is disclosed in each group, see individual claim groupings above for the different sequences. The DNA disclosed in each group is made up of a different sequence having a different structure and different function.

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International application No.

PCT/US01/28861

The special technical feature of group 4, 16 and 31 is considered to be a method of producing recombinant adenoviral particles. Each group contains different sequences hence the resulting particles would have different structures and functions associated with the particle.

The special technical feature of group 5, 17 and 32 is considered to be a method of producing a cellular mediated immune response to the heterologous protein encoded by the different adenoviral vectors. Each group contains different sequences a encoding different protein, therefore the resulting immune response will also be different.

The special technical feature of group 6, 18 and 33 is considered to be a method of producing a cellular mediated immune response to the heterologous protein encoded by the different adenoviral vectors in conjunction with immunizing the individual a DNA plasmid vaccine. Each method contains different sequences encoding a different protein, therefore the resulting immune response will also be different.

Accordingly, groups 1-48 are not so linked by the same or corresponding technical feature as to form a single general inventive concept.

Continuation of B. FIELDS SEARCHED Item 3:

WEST 2.0, STN-BIOSIS, MEDLINE adenoviral vector, deletion, HIV, Gag, polyadenylation signal, CMV promoter

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